

TestAmerica, Inc.

Dayton Division

Quality Assurance Plan

For

Routine Analytical Services

Approved By:

( )

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#### 3. PROJECT DESCRIPTION

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#### 3.1 <u>Introduction</u>

This document describes the essential elements of the Quality Assurance Program at TestAmerica and the quality control procedures utilized by TestAmerica to ensure a national standard of quality.

## 3.2 Scope

TestAmerica believes that quality is the key to maintaining leadership in the environmental analytical industry.

The Quality Assurance (QA) program at TestAmerica requires that each division adhere to a division specific Quality Assurance Plan (QAP) which details the specific quality control procedures for that laboratory; and, as needed, project specific Quality Assurance Project Plans (QAPP).

- 3.2.1 TestAmerica subscribes to the following policies as its standard of quality in its analytical program:
  - It is our policy to maintain a National Quality Assurance program throughout all TestAmerica laboratories, thereby providing our clients with defensible data of known and consistently high quality;
  - It is our policy to communicate the scope and content of our QA Program internally to our employees and to train each employee in the application of our Program;
  - It is our policy that no data will be reported to our clients that has not met our full QA requirements;
  - It is our policy to remove from commercial offering any analysis offered by a TestAmerica laboratory when that laboratory fails to demonstrate that it can consistently perform that analysis to TestAmerica's high quality standard;
  - It is our policy that any employee aware of misrepresentation of facts regarding analytical results is required to notify his/her immediate supervisor or, if this is not feasible, another representative of the management of the company immediately;
  - It is our policy that all personnel be free from any undue internal and external pressures that may adversely affect the quality of their work, including but not necessarily limited to: commercial, client, production, operational and financial influences. Personnel

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believing such pressures exist are required to immediately notify his/her supervisor or, if this is not feasible, another management representative as outlined in the Open Door Policy procedures found in the TestAmerica Human Resource Policy Manual;

- It is our policy to resolve complaints received from clients or other parties regarding the laboratory's activities. The complaint, including when data is questioned, is documented and resolved using the procedures outlined in Section 10. This includes the use of the Inter-Laboratory Notification form and/or the Re-Evaluation Request form. Additionally, the QA Coordinator determines if an audit of the specific activity is necessary;
- It is our policy to notify clients, in writing, when significant doubt is cast on the correctness or validity of data as a result of findings from an audit. The written documentation provides specific findings and conclusions and shall be made using either: the Inter-Laboratory Notification form from Section 10, a letter format, or the content of a Project Case Narrative.

#### 4. PROJECT ORGANIZATION AND RESPONSIBILITY

## 4.1 <u>Introduction</u>

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The success of this QA Plan requires the cooperative efforts and support of all personnel - Divisional and Corporate. The primary responsibility for data quality rests with the analyst in performing frequent and regular quality control checks on the work he/she does. This program is designed to support and coordinate these efforts at the bench level. The divisional organization structure is shown in Figure 4.1 and specific responsibilities related to quality assurance are as follows.

# 4.2 Assignment of Responsibilities

## 4.2.1 The <u>Analyst</u> shall:

- Adhere to analytical and QC protocols prescribed by approved SOPs and QAP/QAPPs;
- Review analytical data and quality control indicators on a daily basis.
- Correct out of control analyses if possible, otherwise, seek the Supervisor's help immediately;
- Meet sample hold times or immediately inform Supervisor if this is not possible;
- Perform routine maintenance on instruments and equipment;
- Maintain all sample tracking, preparation and instrument log books;
- Maintain control charts, as appropriate, to provide real-time trend analysis; - Document out of control situations and their resolution with corrective action reports; and
- Suggest improvements in methodologies to Supervisors and Quality Assurance personnel. These improvements, if approved, will be incorporated into SOPs.

## 4.2.2 The <u>Supervisor</u> (Operations Manager, etc.) shall:

- Provide training for new analysts using approved SOPs, verify adequacy of training and document the training;
- Ensure compliance with approved SOPs and QAP/QAPPs, including the quality control measures they prescribe;

- Investigate and assist the analyst in correcting an out of control analysis, and communicate the corrective action to the Division Manager and the QA Coordinator; Guarantee that sample hold times are met or immediately notify the Division Manager and Customer Service Representative if this cannot be done;
- Assist in the development and revision of SOPs as needed, ensuring that they are: representative of how the procedure is done in the lab, method / technically correct, complete, and of sufficient detail to serve as a training document;
- Monitor control charting maintained by the analysts;
- Review, evaluate and approve data produced by analysts prior to reporting;
- Approve logbook entries for completeness and correctness and ensure that documentation is maintained securely and in an easily retrievable fashion; Assist in the development and revision of the Divisional QAP;
- Serve as a Technical Manager or Deputy Technical Manager if so designated;
- Communicate to the Division Manager any needs for equipment and/or personnel in their area; and
- Communicate with other TestAmerica Supervisors with similar areas of responsibilities.

## 4.2.3 <u>Division Quality Assurance Coordinator</u>

The Division Quality Assurance Coordinator shall:

- Administer the Divisional QA Programs;
- Ensure that a Divisional QA Plan is in place that accurately reflects the QA/QC procedures of the laboratory, and coordinate the revision of the QAP as necessary;
- Assist in the development of SOPs as relates to quality control;
- Serve as the repository for the original copies of SOPs and the QAP and control the distribution of these documents;
- By conducting internal audits, ensure that SOPs are being followed; maintain a list of available SOPs;
- Assist in the writing of QA Project Plans (QAPPs), ensure that they are complete and accurate with regard to

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regulatory requirements, and determine that the laboratory can meet the requirements set forth in the QAPP; maintain a copy of each QAPP;

- Assist in the coordination of PE samples for certification;
- Determine that analysts are properly trained in quality control measures for all analyses;

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- Through internal audits, evaluate quality control processes and documentation throughout the laboratory, making recommendations for improvement when necessary;
- Assist the supervisors and analysts in the use of control charts to monitor analytical performance in the laboratory;
- Assist in interdivisional audits, as appropriate, and serve as QA support to Division Managers in external audits;
- Work closely with the Division Manager to resolve data quality related issues;
- Communicate to the Division Manager areas requiring corrective action and help define appropriate corrective action. Determine that the corrective action has been properly carried out and documented;
- Assist the Division Manager in obtaining and maintaining needed certifications, performance evaluation samples and contract laboratory status;
- Serve as a repository for all audit and performance evaluation results and for certification and licensing documentation;
- Serve as a Technical Manager or Deputy Technical Manager if so designated;
- Communicate with other TestAmerica QA Coordinators; and
- Prepare a monthly QA report and submit to the Division Manager.

# THE DIVISION QUALITY ASSURANCE COORDINATOR WILL NOT:

- Participate in any operational activities involving the production of analytical data or reports. Specifically, his/her responsibilities will not include sample collection, sample receipt or log-in, preparation or analysis of samples, supervision of analytical sections or departments, routine data review, preparation of

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reports, project management, or management of a division.

- Sign analytical reports or data packages to external customers (unless mandated by specific State requirements).

## 4.2.4 The <u>Technical Manager(s)</u> (however named) shall:

- Be designated by the Division Manager;
- Be technically competent in their area of responsibility;
- Have overall technical responsibility for the designated technical operations;
- Provide technical guidance to the analytical staff and be the source point for technical help; and
- Normally hold the position of an Operations Manager or Supervisor but may be a senior analyst in a given department who is readily available to provide technical assistance. There may be more than one Technical Manager, i.e., organic inorganic or departmental, so long as they are properly identified and designated. The Division Manager or QA Coordinator may be a Technical Manager.

# 4.2.5 The <u>Deputy Technical Manager(s)</u> (however named) shall:

- Be nominated by the Division Manager;
- In the temporary absence of the Technical Manager, assume responsibilities for this function;
- Normally hold the position of an Operations Manager (i.e., Inorganic Operations Manager and Organic Operations Manager can serve as each others Deputy), or Supervisor but may be a senior analyst in a given department who is readily available to provide technical assistance. There may be more than one Deputy Technical Manager so long as they are properly identified and designated. The Division Manager or QA Coordinator may be a Deputy Technical Manager.

# 4.2.6 The <u>Division Manager</u> shall:

- In the temporary absence of a Division QA Coordinator, assume all responsibilities of the Division QA Coordinator position;
- Ensure that the operational requirements of this Plan and supporting programs are met;



- Manage the on-going requirements of Quality Assurance and Quality Control activities through Supervisors and Division QA Coordinators;
- Approve and implement SOPs, QAPs and QAPPs;
- Ensure that appropriate corrective actions are taken to address analyses identified as requiring such actions by internal or external performance or procedural audits;
- Review and submit corrective action reports;
- Have in place a system to ensure that sample holding times are met. Notify the client whenever hold times are missed:
- Ensure that all analysts and supervisors have received adequate training to properly carry out the duties assigned to them and document this training;
- Pursue and maintain appropriate laboratory certification and contract approvals. Arrange for the analysis of Performance Evaluation (PE) samples necessary to satisfy certification requirements;
- Serve as a Technical Manager or Deputy Technical Manager if so designated;
- With the help of the Client Service Representative or the Project Manager, ensure that analysts and supervisors know any client specific reporting and QC requirements prior to sample arrival in the lab; and
- Represent, or designate an alternate individual to represent the Division during client and/or regulatory audits, with QA support as needed from Division and/or Corporate QA personnel.

## 4.3 <u>Communications</u>

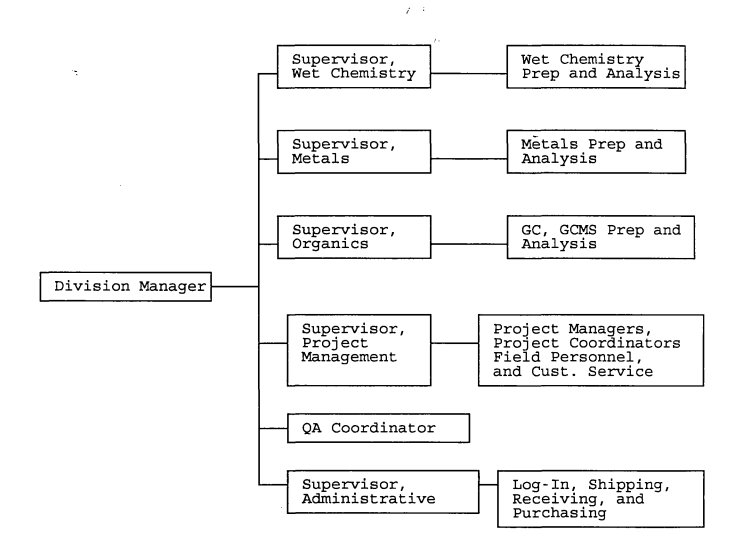
The TestAmerica Corporate office supports an "open door" communications policy: every TestAmerica employee has free access to the Corporate office. Additionally, this Plan supports using resources (people in particular) at all levels; appropriate, frequent, effective communication is encouraged throughout TestAmerica. In addition, specific documents relating to this Plan are available to all employees, including:

Quality Assurance Programs to address specific areas identified in this Plan. Programs exist for SOPs, QAPs, and the Data Quality Audits and these have been incorporated into this Plan.

Quality Assurance Policies to address specific quality related items outside the scope of existing Programs.

Figure 4.1. Organization of TestAmerica, Inc. Dayton Division.

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#### 5. QUALITY ASSURANCE OBJECTIVES

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#### 5.1 Introduction

The quality assurance objectives are to provide analytical data of known and documented quality, to produce defensible analytical data and to produce data which meets the client's specific needs for the data in a cost effective manner.

Data quality is defined in terms of data quality objectives. Data quality objectives are the qualitative and quantitative statements which specify the required data quality based on the end use of the data to be collected. Data quality is assessed by precision, accuracy, representativeness and comparability.

- 5.1.1 To accomplish its data quality objectives, TestAmerica Dayton will:
  - Maintain an effective, on-going Quality Assurance and Quality Control Program that measures and verifies laboratory performance;
  - Provide sufficient flexibility to allow controlled changes in routine methodology to meet project specific data requirements;
  - Recognize as soon as possible and provide correction for any factors which may adversely affect data quality;
  - Monitor operational performance of the laboratory on a routine basis and provide corrective action as needed;
  - Maintain complete records of sample submittal, raw data, laboratory performance and complete analysis to support reported data.

# 5.2 <u>Level of Quality Control and Quality Assurance Efforts</u>

TestAmerica maintains a well defined internal quality control (QC) program. A system of specific activities are in use in the laboratory to ensure that the analytical data generated is of consistently high quality. Blanks, Calibration Verification Standards, Laboratory Control Samples, Spikes, Duplicates and Matrix Spikes are analyzed and monitored at regular frequencies, to ensure that the data quality objectives for the project are met.

#### 5.3 Accuracy

Accuracy is defined as how close an analytical value is to the actual concentration of analyte in the sample. Accuracy is evaluated through the analysis of Laboratory Control Samples

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(LCS). Matrix Spikes may also be used to assess accuracy. Accuracy goals are outlined in Table 5.1 through 5.11.

#### 5.4 Precision

Precision is defined as the repeatability of a measurement. It is an indication of the variability of a measurement. Precision is evaluated through the use of matrix spike/matrix spike duplicates (MS/MSD) or through duplicate analysis when matrix spiking is not possible. Precision is expressed in terms of relative percent difference (RPD). Precision goals are outlined in Table 5.1 through 5.11.

# 5.5 <u>Completeness</u>

Completeness is defined as the measure of the amount of valid data, as determined utilizing the quality assurance and the associated standard operating procedures, obtained from the analytical measurement system compared to the amount of valid data that was expected to be obtained under correct operating conditions. Completeness is expressed as a percentage of the number of data with acceptable results divided by the number of expected results.

Completeness will be determined by the client. Ideally, all of the analyses will be valid. However, some samples may be lost in laboratory accidents or some results may be deemed questionable based on internal quality control. TestAmerica will make every effort to produce analytical data that meets the completeness requirements of the client.

#### 5.6 Representativeness

Representativeness is a measure of how closely the analytical results reflect the actual concentration of analytes in the sample. For any project, sampling will be performed by the customer or the customer's representative (the customer may contract with TestAmerica for sampling services). Sample handling protocols (i.e., storage and preservation) have been developed to preserve the representativeness of the collected samples.

Every attempt will be made to ensure that the aliquots taken for analysis are representative of the sample received. TestAmerica will notify the client if samples received in the laboratory have any of the following conditions: improper preservation, broken sample containers, chain of custody discrepancies, broken or missing custody seals (if required) and TestAmerica will document such deviations. All other measures of representativeness will be determined by the client.

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# <u>5.7</u> Comparability

The generation of comparable data is the goal of any analytical program. This characteristic implies strict adherence to published analytical protocols and use of standard reporting units. TestAmerica's QC program is structured to ensure adherence to the proper analysis protocols and fully document these procedures. The QA objective is that all data resulting from these analyses be comparable with other measurements made by TestAmerica or another organization. All judgements of comparability will be made by the client.

## 5.8 Quality Control Measures

The following tables summarize the Quality Control Indicators (QCIs) which are performed with the common analytical procedures at TestAmerica-Dayton. The tables are for general reference, as method specific criteria varies. Please refer to the Standard Operating Procedures (SOPs) for specific control limit information.

Table 5.1. Quality Control Measures for Wet Chemistry

Quality Control Measure	Frequency	Control Limits
Calibration Curve	*	Correlation Coef. ≥ 0.995
Initial Calibration Verification (ICV)	1 / Calibration	Accuracy 90 - 110 %
Reagent Blank	Daily	< Reporting Limit
Method Blank	1 / 20 samples	< Reporting Limit
Continuing Calibration Verification (CCV)	Beginning & end of run; 1 / 10 samples	**
Laboratory Control Sample (LCS)	1 / batch	**
Matrix Spike/ Matrix Spike Duplicate (MS/MSD)	1 / batch	**
Duplicate	1 / batch if parameters cannot be spiked	**

<sup>\*</sup> If calibrations are applicable to a Wet Chemistry parameter, they will be performed on a daily basis, or at the frequency specified in the SOP.

<sup>\*\*</sup> The control limits for these Quality Control Indicators are statistically determined annually based on +/- 3 standard deviations from the mean. Control limits can not exceed the range listed in the method.

Table 5.2. Quality Control Measures for Bacterial Analyses
Fecal Coliform

Quality Control Measure	Frequency	Control Limits
Media pH Control	Weekly	+/- 0.2 pH Units
Filtration Blanks	Daily	< 1 Colony

#### Total Coliform

Quality Control Measure	Frequency	Control Limits	
Media Quality Check	1/ Media Batch	E. Coli + Klebsiella + Pseudomonas -	
Positive control	1/ Sample Set	Positive Coliform	
Negative control	1/ Sample Set	Negative Coliform	
Sample bottle sterility check	3/ box of sample bottles	Negative Coliform	

Table 5.3. Quality Control Measures for Metals Graphite Furnace

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Quality Control Measure	Frequency // :	Control Limits
Calibration Curve	Daily	Correlation Coef. ≥ 0.995
Initial Calibration Verification (ICV)	Daily	Accuracy 90 - 110 %
Reagent Blank	Daily	< Reporting Limit
Method Blank	1 / Batch	< Reporting Limit
Continuing Calibration Verification	Beginning & end of run 1 / 10 samples	Accuracy 90 - 110 %
Laboratory Control Sample (LCS)	1 / Batch	80 % - 120 %
Matrix Spike/ Matrix Spike Duplicate (MS/MSD)	1 / Batch	75 % - 125 %

Table 5.4. Quality Control Measures for Metals by Inductively Coupled Plasma Atomic Emission (ICP-AES)

*"*:

Quality Control Measure	Frequency	Control Limits		
Calibration Curve	Daily	% RSD of three readings		
Re-analyze Calibration Standards	Daily	Accuracy 95 - 105 %		
Initial Calibration Verification (ICV)	Daily	Accuracy 90 - 110 %		
Reagent Blank	1 / 10 Samples	< Reporting Limit		
Reporting Limit Verification (RLV)	Daily	Accuracy 70 - 130 %		
Spectral Interference Checks (SIC)	Beginning & end of run	Per Method**		
Continuing Calibration Verification (CCV)	Beginning & end of run; 1 / 10 Samples	Accuracy 90 - 110 %		
Method Blanks	1 / Batch	< Reporting Limit		
Laboratory Control Samples (LCS)	1 / Batch	85 % - 115 %		
Matrix Spike/ Matrix Spike Duplicates	1 / Batch	75 % - 125 %		

<sup>\*\*</sup> Please refer to the SOP for Method specific criteria.

Table 5.5. Quality Control Measures for Metals by Inductively Coupled Plasma-Mass Spectrometry (ICP-MS)

# US EPA Method 200.8

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Quality Control Measure	Frequency	Control Limits
Calibration Blank	Daily	
Calibration Curve	Daily	
Initial Calibration Verification (ICV)	Daily	- Accuracy 90 - 110 %
Continuing Calibration Blank	Beginning & end of run 1 / 10 Samples	< 1/10 Reporting Limit or 2.2x the MDL, which ever is greater
Reporting Limit Verification (RLV)	Daily	Accuracy 70 - 130 %
Continuing Calibration Verification (CCV)	Beginning & end of run; 1 / 10 Samples	Accuracy 90 - 110 %
Reagent Blank	1 / Batch	< 1/10 Reporting Limit or 2.2x the MDL, which ever is greater
Laboratory Control Samples (LCS)	1 / Batch	85 % - 115 %
Matrix Spike/ Matrix Spike Duplicates (MS/MSD)	1 pair / Batch	75 % - 125 %
Internal Standard	All	Accuracy 60 - 125 % of Initial Cal. Blank
Mass Calibration and Resolution Check	Daily	Per Method**
Instrument Stability	Daily	Per Method**

NOTE: Rinse Blanks are used after each Quality Control or client sample.

<sup>\*\*</sup> Please refer to the SOP for Method specific criteria.

Table 5.6. Quality Control Measures for Metals by Inductively Coupled Plasma-Mass Spectrometry (ICP-MS)

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#### SW 846 Method 6020

Quality Control Measure	Frequency	Control Limits
Calibration Blank	Daily	
Calibration Curve	Daily	
Initial Calibration Verification (ICV)	Daily	Accuracy 90 - 110 %
Continuing Calibration Blank	Beginning & end of ru 1 / 10 Samples	n; < Reporting Limit
Reporting Limit Verification (RLV)	Daily	Accuracy 70 - 130 %
Continuing Calibration Verification (CCV)	Beginning & end of ru 1 / 10 Samples	n; Accuracy 90 - 110 %
Reagent Blank	1 / Batch	< Reporting Limit
Laboratory Control Samples (LCS)	1 / Batch	, 85 % - 115 %
Matrix Spike/ Matrix Spike Duplicates (MS/MSD)	1 / Batch	75 % - 125 %
Internal Standard	Initia - Accu Initia	aracy 30 - 120 % of al Cal. Blank for samples aracy 80 - 120 % of al Cal. Blank for Quality ol samples
Mass Calibration and Resolution Check	Daily	Per Method**
Instrument Stability	Daily	Per Method**
Interference Check Sample	Beginning & end of ru	n Per Method**

NOTE: Rinse blanks are used after each Quality Control or client sample.

<sup>\*\*</sup> Please refer to the SOP for Method specific criteria.

Table 5.7. Quality Control Measures for Mercury by Cold Vapor

Quality Control Measure	Frequency /	Control Limits	
Calibration Curve	Daily	Correlation Coef. ≥ 0.995	
Initial Calibration Verification (ICV)	Daily	Accuracy 90 - 110	
Reagent Blank	Daily	< Reporting Limit	
Method Blank	1 / Batch	< Reporting Limit	
Continuing Calibration Verification	Beginning & end of run 1 / 10 samples	Accuracy 80 - 120	
Matrix Spike/ Matrix Spike Duplicate (MS/MSD)	1 / Batch	75 % - 125 %	

Table 5.8. Quality Control Measures for Volatiles by GC/MS

#### US EPA Method 624

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Quality Control Measure	Frequency	Control Limits
Initial Calibration	*	Minimum of 3 Standards < 35% RSD
Initial Calibration Verification (ICV)	1 / Calibration	± 30 % of True Value
Tune Check	1 / 12 hours	Per Method**
Continuing Calibration Verification (CCV)	1 / 12 hours	Per Method**
Surrogates/Internal Standards	All	Per Method**
Method Blanks	1 / 12 hours	< Reporting Limit
Matrix Spike/ Laboratory Control Standards	1 / 20 samples	Per Method**

<sup>\*</sup> An initial calibration is required whenever the Quality Control Indicators do not pass established acceptance criteria.

\*\* Please refer to the SOP for Method specific criteria.

# SW 846 8260A

Quality Control Measure	Frequency	Control Limits
Initial Calibration	*	Minimum of 5 Standards SPCC/CCC per Method**
Initial Calibration Verification (ICV)	1 / Calibration	± 30 % of True Value
Tune Check	1 / 12 hours	Per Method**
Continuing Calibration Verification (CCV)	1 / 12 hours	SPCC/CCC per Method**
Surrogates/Internal Standards	All	Per Method**
Method Blanks	1 / 12 hours	< Reporting Limit
Matrix Spike/ Matrix Spike Duplicate & Laboratory Control Standard	1 / 20 Samples and/or daily	Per Method**

Table 5.8. Continued...

#### US EPA Method 524.2

Quality Control Measure	Frequency	Control Limits
Initial Calibration	*	Minimum of 4 standards
Initial Calibration Verification (ICV)	1 / Calibration	± 40 % of True Value
Tune Check	1 / 12 hours	Per Method**
Reporting Limit Verification Standard	1 / 12 hours	± 40 % of True Value
Continuing Calibration Verification (CCV)	1 / 12 hours up to 20 samples	< 30 % RSD
Surrogates/Internal Standards	All	Per Method**
Reagent Blank	1 / 12 Hours	< Reporting Limit

<sup>\*</sup> An initial calibration is required whenever the Quality Control Indicators do not pass established acceptance criteria.

\*\* Please refer to the SOP for Method specific criteria.

Table 5.9. Quality Control Measures for Semi-volatiles by GC/MS
SW 846 8270B

Quality Control Measure	Frequency	Control Limits
Initial Calibration	*	Minimum of 5 Standards < 30 % RSD
Initial Calibration Verification (ICV)	1 / Calibration	± 30 % of True Value
Tune Check	1 / 12 hours	Per Method**
Continuing Calibration Verification (CCV)	1 / 12 hours	SPCC/CCC per Method**
Surrogates/Internal Stand	ards All	Per Method**
Method Blanks	1 / Extraction Set	< Reporting Limit
Matrix Spike/ Matrix Spike Duplicate	1 / 20 Samples	Per Method**
Laboratory Control Standard	1 / Extraction Set	Per Method**
	US EPA Method 625	
Quality Control Measure	Frequency	Control Limits
Initial Calibration	*	Minimum of 3 Standards < 35 % RSD
Initial Calibration Verification (ICV)	1 / Calibration	+ 30 % of True Value
Tune Check	1 / 12 hours	Per Method**
Continuing Calibration Verification (CCV)	1 / 12 hours	Per Method**
Surrogates/Internal Stand	ards All	Per Method**
Method Blanks	1 / Extraction Set	< Reporting Limit
Laboratory Control Standard	1 / Extraction Set	Per Method**
Matrix Spike/ Matrix Spike Duplicate	1 / 20 Samples	Per Method**

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# Table 5.9. Continued...

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\* An intial calibration is required whenever the Quality Control Indicators do not pass established acceptance criteria.

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\*\* Please refer to the SOP for Method specific criteria.

Table 5.10. Quality Control Measures for Pesticides/PCBs - ----SW 846 8080A

Quality Control Measure	Frequency	Control Limits	
Initial Calibration	*	Minimum of 5 Standards < 20 % RSD	
Initial Calibration Verification (ICV)	1 / Calibration	± 30 % of True Value	
Continuing Calibration Verification (CCV)	Beginning & end of run; 1 / 10 samples	Per Method**	
Method Blank	1 / Extraction Set	< Reporting Limit	
Surrogates	All	Per Method**	
Laboratory Control Sample (LCS)	1 / Extraction Set	Per Method**	
Matrix Spike/ Matrix Spike Duplicate	1 / 20 Samples	Per Method**	
	US EPA Method 608		
Quality Control Measure	Frequency	Control Limits	
Initial Calibration	*	Minimum of 3 Standards < 10 % RSD	
Initial Calibration Verification (ICV)	1 / Calibration	± 30 % of True Value	
Continuing Calibration Verification (CCV)	Beginning & end of run; 1 / 10 samples	< 15 % Difference	
Method Blank	1 / Extraction Set	< Reporting Limit	
Surrogates	All	Per Method**	
Laboratory Control Sample (LCS)	1 / Extraction Set	Per Method**	
Matrix Spike/ Matrix Spike Duplicate	1 / 20 Samples	Per Method**	

<sup>\*</sup> An initial calibration is required whenever the Quality Control Indicators do not pass established acceptance criteria.

\*\* Please refer to the SOP for Method specific criteria.

Table 5.11. Quality Control Measures for Total Petroleum Hydrocarbons (TPH) by FTIR

## US EPA Method 418.1

Quality Control Measure	Frequency	Control Limits  Min. of 3 standards Correlation Coef. ≥ 0.995  Accuracy 90 - 110%	
Initial Calibration	*, Daily for Voluntary Action Program		
Initial Calibration Verification (ICV)	1 / Calibration		
Reagent Blank	Daily	< Reporting Limit	
Method Blank	1 / Extraction Set	< Reporting Limit	
Continuing Calibration Verification (CCV)	1 / 10 Samples	**	
Laboratory Control Samples (LCS)	1 / Extraction Set	**	
Matrix Spike/ Matrix Spike Duplicate	1 / 20 Samples	**	

<sup>\*</sup> An initial calibration is required whenever the Quality Control Indicators do not pass established acceptance criteria.

<sup>\*\*</sup> The control limits for these Quality Control Indicators are statistically determined annually based on +/- 3 standard deviations from the mean. Control limits can not exceed the range listed in the method.

Table 5.12. Quality Control Measures for Total Petroleum Hydrocarbons (Diesel Range Organics)

# SW-846 8015B

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Quality Control Measure	Frequency	Control Limits  Min. of 5 standards % RSD < 20% or Correlation Coef. ≥-0.995	
Initial Calibration	*		
Reagent Blank	Daily	< Reporting Limit	
Method Blank	1 / Extraction Set	< Reporting Limit	
Continuing Calibration Verification (CCV)	1 / 20 Samples up to 12 hours	< 15 % Difference	
Laboratory Control Samples (LCS)	1 / Extraction Set	**	
Matrix Spike/ Matrix Spike Duplicate	1 / 20 Samples	**	
Surrogate	1 / Sample	**	

\* An initial calibration is required whenever the Quality Control Indicators do not pass established acceptance criteria.

<sup>\*\*</sup> The control limits for these Quality Control Indicators are statistically determined annually based on +/- 3 standard deviations from the mean. Control limits can not exceed the range listed in the method.

Quality Control Measures for Total Petroleum Hydrocarbons Table\_5.13. (Gasoline Range Organics)

#### SW-846 8015A Modified

% RSD < 20% of Correlation of ≥ -0.99  Reagent Blank  Daily  Continuing Calibration  1 / 10 Samples  < 15 % Differ Verification (CCV)  Laboratory Control  1 / Batch up to		·
% RSD < 20% of Correlation of ≥ -0.99  Reagent Blank  Daily  Continuing Calibration  Verification (CCV)  Laboratory Control  Patch up to		As = = = = = = = = = = = = = = = = = = =
Continuing Calibration 1 / 10 Samples < 15 % Differ Verification (CCV)  Laboratory Control 1 / Batch up to	tial Calibration	Min. of 5 standard % RSD < 20% or Correlation Coef. ≥-0.99
Verification (CCV)  Laboratory Control 1 / Batch up to	gent Blank	< Reporting Limit
		mples < 15 % Difference
samples (LCS)	oratory Control ples (LCS)	
Matrix Spike/ 1 pair / Batch up Matrix Spike Duplicate to 20 samples **		
Surrogates All Samples/Standards **	rogates	es/Standards **

<sup>\*</sup> An initial calibration is required whenever the Quality Control

Indicators do not pass established acceptance criteria.

\*\* The control limits for these Quality Control Indicators are statistically determined annually based on +/- 3 standard deviations from the mean. Control limits can not exceed the range listed in the method.

Quality Control Measures for Total Petroleum Hydrocarbons Table 5.14. (Gasoline Range Organics)

#### SW-846 8015B/ +

Quality Control Measure	Frequency	Control Limits	
*			
Initial Calibration	*	Min. of 5 standards % RSD < 20% or Correlation Coef. ≥-0.99	
Reagent Blank	Daily	< Reporting Limit	
Continuing Calibration Verification (CCV)	1 / 20 Samples up to 12 hours	< 15 % Difference	
Laboratory Control Samples (LCS)	1 / Batch up to 20 samples	**	
Matrix Spike/ Matrix Spike Duplicate	1 pair / Batch up to 20 samples	**	
Surrogates	All Samples/Standards	**	

<sup>\*</sup> An initial calibration is required whenever the Quality Control Indicators do not pass established acceptance criteria.

\*\* The control limits for these Quality Control Indicators are statistically determined annually based on +/- 3 standard deviations from the mean. Control limits can not exceed the range listed in the method.

Table 5.15. Quality Control Measures for BTEX SW-846 8020A

Quality Control Measure	Frequency	Control Limits	
Initial Calibration	*	Min. of 5 standards % RSD < 20% or Correlation Coef. ≥ 0.99	
Reagent Blank	Daily	< Reporting Limit	
Continuing Calibration Verification (CCV)	1 / 10 Samples	< 15 % Difference	
Laboratory Control Samples (LCS)	1 / Batch up to 20 samples	· **	
Matrix Spike/ Matrix Spike Duplicate	1 pair / Batch up to 20 samples	**	
Surrogates	All Samples/Standards	; **	

<sup>\*</sup> An initial calibration is required whenever the Quality Control

Indicators do not pass established acceptance criteria.

\*\* The control limits for these Quality Control Indicators are statistically determined annually based on +/- 3 standard deviations from the mean. Control limits can not exceed the range listed in the method.

Table 5.16. Quality Control Measures for BTEX SW-846 8021B

Quality Control Measure	Frequency	Control Limits  Min. of 5 standards % RSD < 20% or Correlation Coef. ≥ 0.99	
Initial Calibration	*		
Reagent Blank	Daily	< Reporting Limit	
Continuing Calibration Verification (CCV)	1 / 20 Samples up to 12 hours	< 15 % Difference	
Laboratory Control Samples (LCS)	1 / Batch up to 20 samples	**	
Matrix Spike/ Matrix Spike Duplicate	1 pair / Batch up to 20 samples	**	
Surrogates	All Samples/Standards	**	

<sup>\*</sup> An initial calibration is required whenever the Quality Control Indicators do not pass established acceptance criteria.

\*\* The control limits for these Quality Control Indicators are statistically determined annually based on +/- 3 standard deviations from the mean. Control limits can not exceed the range listed in the method.

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Table 5.17. Polynuclear Aromatic Hydrocarbons by HPLC

SW-846 8310

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Quality Control Measure	Frequency	Control Limits  Min. of 5 standards % RSD < 20% or Correlation Coef. ≥ 0.99	
Initial Calibration	*		
Reagent Blank	Daily	< Reporting Limit	
Method Blank	1 / Extraction Set	< Reporting Limit	
Continuing Calibration Verification (CCV)	1 / 20 Samples up to 12 hours	< 15 % Difference	
Laboratory Control Samples (LCS)	1 / Extraction Set	**	
Matrix Spike/ Matrix Spike Duplicate	1 / 20 Samples	**	
Surrogate	1 / Sample	**	

<sup>\*</sup> An initial calibration is required whenever the Quality Control Indicators do not pass established acceptance criteria.

\*\* The control limits for these Quality Control Indicators are statistically determined annually based on +/- 3 standard deviations from the mean. Control limits can not exceed the range listed in the method.

Table 5.18. Quality Control Measures for Radiological Parameters

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· CROSS			
Quality Control Measure	Frequency	Control Limits	
Calibration of Efficiency Factor	Annually	NA	
Method Blank	1 / 20 Samples	< Reporting Limit	
Continuing Calibration Verification (CCV)	Daily	<u>.</u> *	
Laboratory Control Samples (LCS)	1 / 20 Samples	*	
Matrix Spike / Matrix Spike Duplicates	1 / 20 Samples	. *	

<sup>\*</sup> The control limits for these Quality Control Indicators are statistically determined annually based on +/- 3 standard deviations from the mean.

#### 6. SAMPLING PROCEDURES

Often, field sampling is the most critical aspect of an analysis. To ensure the reliability of the data, quality control measures are included in all field sampling activities completed by TestAmerica personnel. Result validity, is aided by proper sampling, handling and identification of samples through detailed chain-of-custody procedures.

#### 6.1 Sampling

The sampling site is chosen by the client. Sampling points are documented as to their exact location for purposes of future sampling.

TestAmerica provides sample media, containers and preservatives as outlined in Table 6.1, as well as shipping containers (coolers) for any project accepted by TestAmerica. A chain of custody record will be provided with each set of sample containers supplied. Chain of custody records are described in more detail in Section 7 of this document.

All field sampling equipment used by TestAmerica is thoroughly cleaned with lab detergent and water and a stiff brush. Field sampling equipment is decontaminated between samples in the field.

When sampling is performed by TestAmerica, background information is gathered to determine if any safety risks are involved in sampling. This background information is also used to make decisions on what type of sampler to use, type of sample container to use and number of samples to take.

#### 6.2 <u>Sample Types</u>

6.2.1 The two most common types of field samples are the grab sample and the composite sample. The definitions of grab and composite samples are as follows:

**Grab** A discrete aliquot that is representative of one specific sample site, at a specific point in time. The entire sample is collected at one point and all at one time.

**Composite** A sample composed of more than one specific aliquot collected at various sites and/or at different points in time.

6.2.2 Blanks can also be collected during the sampling process. The three main types of blanks associated with sampling are the field blank, the trip blank and the equipment blank.

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The definitions of the various types of blanks are as follows:

Field Blank A field blank is an aliquot of analyte-free water that is brought to the field site in a sealed sample container, poured into the appropriate sample containers and transported back to the laboratory. Field blanks are used to determine previously existing container or preservative contamination, and/or contamination that may have resulted from existing field conditions when samples were collected.

Equipment Blank A sample of analyte free water that is poured appropriately over or through the sampling device, containerized, preserved (if the samples are preserved) and handled in the same manner as the samples. The equipment blank is used to identify sample contamination (if any) acquired through collection, handling, preservation and transport.

Trip Blank A sample of analyte-free water which is taken before the sampling event has begun. The trip blank travels with the sample containers as they are shipped to the field site and as the samples are sent back to the laboratory. The trip blank is not opened in the field. It is used to identify contamination or cross contamination due to location or shipping conditions.

All sample types should be maintained during shipment at 4 degrees Celsius. Table 6.1 lists common sample containers and preservatives.

#### 6.3 <u>Subcontracted Analyses</u>

The laboratory will endeavor to inform clients prior to subcontracting analyses to other laboratories. When this subcontracting is routine, the client will be informed by letter or by notation on the sample bottle order included in all bottle shipments. Data from subcontracted analyses are flagged on the analytical data reports.

Table 6.1. General Guidelines for Samples

The contract datacrass for samples				
Parameter	Container	Preservative	Volume	Hold Time
		J. 1		
General Chemistry		<b>.</b>		
Acidity	P,G	4°C	100	14 days
Alkalinity	P,G	4°C	100	14 days
BOD/CBOD	P,G	4°C	1000	48 hours
Chloride	P,G	None	100	28 days
Chlorine	P,G	None	200	On site
COD	P,G	4°C,H2SO4, pH <2	100	28 days
Color	P,G	4°C	50	48 hours
Cyanide, Amenable	P,G	4°C,NaOH, pH >12	1000	14 days
Cyanide, Total	P,G	4°C,NaOH, pH >12	1000	14 days
Fluoride, Total	P	None	300	28 days
Hardness	P,G	4°C,HNO3, pH <2	100	6 months
Ignitability	G	None	100	
Nitrogen, Ammonia	P,G	4°C,H2SO4, pH <2	400	28 days
Nitrogen, Kjeldahl	P,G	4°C,H2SO4, pH<2	500	28 days
Nitrogen, Nitrite	P,G	4°C	100	48 hours
Nitrogen, Nitrate & Nitrit	ce P,G	4°C,H2SO4, pH <2	100	28 days

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Table 6.1 Continued...

Parameter	Container	Preservative	Volume	Hold Time
Oil & Grease	G .	4°С,Н2SO4, рН <2	1000	28 days
Paint Filter, Liquids	G	None	250	NA
рн	P,G	None	25	On Site
Phenols	G	4°C,H2SO4, pH <2	500	28 days
Phosphorus, Ortho	P,G	4°C	100	48 hours
Phosphorus, Total	P,G	4°C,H2SO4, pH <2	100	28 days
Residue, Filterable (TDS)	P,G	4°C	500	7 days
Residue, Non-Filterable (TSS	S) P,G	4°C	500	7 days
Residue, Settleable (SS)	P,G	4°C	1000	7 days
Residue, Total (TS)	P,G	4°C	500	7 days
Residue, Volatile (TVS)	P,G	4°C	500	7 days
Specific Conductance	P,G	4°C	100	28 days
Sulfate	P,G	4°C	100	28 days
Sulfide	P,G	4°C,NaOH,ZnAc pH >9	, 500	7 days
Sulfite	P,G	None	100	on site
Surfactants (MBAS)	P,G	4°C	250	48 hours
Total Organic Carbon (TOC)	P,G	4°C,H2SO4, pH <2	250	28 days
Turbidity	P,G	4°C	100	2 days
Bacteria				
Coliform, Fecal	P ster	ile None	100	6 hours
Coliform, Total and E. Coli	P ster	ile None	100	30 hours

Table 6.1 Continued...

Table 6.1 Continued				
Parameter	Container	Preservative	Volume	Hold Time
<b>Metals</b>		./		
Chromium, Hexavalent	P	4°C	500	24 Hours
Mercury	P	HNO3, pH <2	250	28 Days
All Other Metals	P	HNO3, pH <2	250	6 months
TCLP	G	4°C	1000 g	14 days
Radiological			_	
Alpha/Beta	P	$HNO_3$ , $pH < 2$	1000	6 months
Organics				
Volatile Organics **	G Vials	4°C,HCl, pH <2 *	40 (x3)	14 days
Pesticides/PCB's	G	4°C *	1000	7 days
Pesticides	G	4°C *	1000	7 days
Extractable Organics	G	4°C *	1000	7 days
PNAs	G	4°C *	1000	7 days
TPH (418.1, DRO)	G	4°C,HCl, pH <2 *	1000	7 days
TPH (GRO) **	G	4°C,HCl, pH <2 *	40 (x3)	14 days
Petroleum Hydrocarbons	G	4°C	1000	7 days
Hydrocarbon Solvents	G	4°C	25	NA
PCB in Oils	G Vial	None	2	NA
TCLP	G	4°C	1000 g	14 days

Solids and soils are collected in wide mouth glass jars which have Teflon-lined lids. Samples are maintained at 4°C, if required.

<sup>\*</sup> NOTE: Chlorinated water sources must first be dechlorinated.

<sup>\*\*</sup> Soil samples for SW-5035 are collected in triplicate with Encore samplers and preserved at the laboratory within 48 hours with sodium bisulfate and/or methanol. If field preservation is required, two vials with sodium bisulfate and one vial with 5 mL of Methanol are provided for collecting soil samples.

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#### 7. SAMPLE CUSTODY

Correct sample handling procedures are an integral part of the Quality Assurance program for TestAmerica. A chain of custody documents the sample identity, number of samples, requested analyses and the custody of samples.

# 7.1 Chain of Custody Procedures

Chain of Custody forms are utilized to document, in a legally defensible manner, the transfer of custody for each sample. TestAmerica will follow the descriptions and requested analyses outlined on the Chain of Custody provided by the client. TestAmerica strongly recommends that the chain of custody (COC) be completed and sent with the samples to the lab for analysis. Failure to submit a COC may result in delays for laboratory analysis and possible legal problems if the site evaluation comes into question at a later date.

When samples arrive at TestAmerica, the Sample Custodian documents the condition of custody seals on the Chain of Custody. The temperature of the cooler is documented. The sample custodian checks the sample label against the chain of custody, and notes any deviations. In cases where there are discrepancies between the samples received and the COC, or when samples are received damaged, incorrectly preserved or missing, TestAmerica will notify the client and require that any changes be submitted to TestAmerica in writing.

Samples are then logged into TestAmerica's Laboratory Information Management System (LIMS) and are assigned a unique sample identification number and the requested analyses are linked to the identification number.

Samples that require temperature preservation are maintained at approximately 4 degrees Celsius in a designated sample storage area until the time of analysis and are returned to this area when not in the custody of an analyst.

## 7.2 <u>Laboratory Document Control</u>

All documentation in logbooks and other pertinent documents are entered in ink. Corrections made to data are performed in accordance with EPA Guidelines.

All raw data and pertinent records are maintained for a period of 7 years for non-potable data and 10 years for potable data. As part of the Voluntary Action Program (VAP) requirements, all documents prepared or acquired in connection with a voluntary action will be retained for a period of ten years from the date the analyses were submitted to a certified professional.

#### 8. CALIBRATION PROCEDURES AND FREQUENCY

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This section describes the calibration procedures and frequency for the instrumentation which will be used in the determination of the parameters of interest.

# 8.1 <u>Laboratory Standards</u>

All materials used for instrument calibration, internal standards and surrogate standards will be of the highest purity available from a commercial source. All standards will have a minimum purity of 96%. The calibration procedures outlined here are those routinely used in the laboratory. The calibration frequencies are listed in the Tables in Section 5.

# 8.2 <u>Standards Traceability</u>

All materials, whether high purity bulk material or prepared solutions, will have the following information, at a minimum, recorded into an analytical standards logbook: identity, supplier, lot number, date received, reported concentration and expiration data. This information will be recorded when the material is received or no later than the first time the material is opened.

All analytical standards and spiking solutions will have a unique identification consisting of a name, concentration, expiration date, logbook reference number and the preparation or received date. This identification will be clearly recorded on the label of any bottle containing this material. By consistently using this identification on raw data, the solution can be traced back to the original material.

Documentation of all standard preparations will be recorded in logbooks. The volume and numerical reference of all analytical standards or spiking solutions used in the preparation of another standard will be recorded in the standard preparation logbook.

All calibration standards must be verified against an independently prepared standard from a second manufacturer or a different lot from the same manufacturer.

#### 8.3 Instrument Calibration

Instrument calibration is described in detail in the method specific Standard Operating Procedures. Please refer to the SOPS for additional information concerning calibration and the associated Quality Control Indicators.

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# 8.4. Analytical Balances

Analytical balance calibration is verified on a monthly and daily basis with NBS traceable class S weights. The calibration of each analytical balance is checked on a daily basis by the use of two weights, one in the milligram range and one in the gram range, to determine if the calibration is still valid. A more thorough validation is done on a monthly basis with four weights. All analytical balances receive yearly system checks and calibrations from certified technicians.

# 8.5. Non-analytical Laboratory Equipment

Laboratory equipment, such as ovens and refrigerators which are required to maintain specific temperature ranges, will be monitored daily with thermometers that are calibrated annually against an NIST certified thermometer. For oven temperature requirements, please refer to the method specific SOPs. Freezer temperatures must be maintained between -10°C and -20°C. The refrigerator must be maintained at 4°C.

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# 9. ANALYTICAL PROCEDURES

The Dayton Division of TestAmerica Inc. uses a wide range of analytical methodology for the analysis of wastewater, groundwater, drinking water, and hazardous waste. The tables in this Section list the methods routinely performed.

# 9.1 Methodology

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The analytical methodology performed by TestAmerica conforms to acceptable methods as listed in the governing environmental regulations. Methods are referenced from Standard-Methods for the Examination of Water and Wastewater; U.S. EPA Manual 600/4-79-020, "Methods of Chemical Analysis of Water and Wastes"; U.S. EPA Manual SW-846, Test Methods for Evaluating Solid Waste"; relevant ASTM, NIOSH and other publications.

The methods listed in Tables 9.1 through 9.5 are representative of analyses which are routinely performed. This laboratory has the capability to perform other methods. If a method of interest is not listed in this document, consult a Customer Service Representative or Project Manager to see if the laboratory is capable of performing the analysis.

## 9.2 Reporting Limits

TestAmerica has established reporting limits for all routine analyses. Ideally, reporting limits are based on the Limit of Quantitation (LOQ) that was determined when method detection limit studies were performed. Due to permit requirements or other client requirements it may be necessary to report at a value below the LOQ but still above the MDL. At no time will results be reported at less than the calculated MDL. The LOQ is defined as the level above which quantitative results may be obtained with a specified degree of confidence. The LOQ is calculated as ten times the standard deviation of the population of data obtained in the method detection limit study.

Method detection limit studies are performed annually on all analytes. These studies are performed in accordance with procedures in CFR Part 136 Appendix B.

The tables 1 show the reporting limits used by TestAmerica Dayton. Reporting limits listed are based on minimal matrix interference for aqueous samples. Actual reporting limits may vary due to sample matrix and sample dilution requirements.

Table 9.1. Analytical Methods and Reporting Limits - Potables

Parameter	Method Reference	Method Description	Report Limit	ing
Vet Chemistry			<u></u>	
Alkalinity	SM 2320	Titration	10	mg/L
Chloride	SM 4500Cl-B	Argentometric	` 5	mg/L
Total Residual Chlorine	SM 4500Cl-G	DPD Colorimetric ~	0.1	mg/L
Coliform, Total	MMO-MUG	Colilert/Colisure	Preser	nce/Absence
Coliform, E. Coli	MMO-MUG	Colilert/Colisure	Preser	nce/Absence
Cyanide, Total	EPA-335.4	Spectrophotometric	0.005	mg/L
Fluoride	SM 4500F-C	Ion-Selective Electrode	0.05	mg/L
Gross Alpha	EPA 900.0	Alpha Emission	3	pCi/L
Gross Beta	EPA 900.0	Beta Emission	4	pCi/L
Hardness, Total (CaCO3)	EPA-130.2	Titration, EDTA	5	mg/L
Nitrogen, Nitrate	SM 4500NO3-F	Automated Cd Reduction	0.02	mg/L
Nitrogen, Nitrite	SM 4500NO3-F	Automated Cd Reduction	0.02	mg/L
Nitrogen, Nitrate + Nitrite	SM 4500NO3-F	Automated Cd Reduction	0.02	mg/L
рН	EPA-150.1	Potentiometric	0.1	s.u.
Phosphorus, Total	SM 4500P-E	Spectrophotometric	0.10	mg/L
Stability	SM 2330	Calcium Carbonate Saturation	NA	
Total Dissolved Solids	SM 2540 C	Gravimetric, 180°C	50	mg/L
Turbidity	EPA-180.1	Nephelometric	1.0	NTU
detals				
Aluminum (Al)	EPA-200.7 EPA-200.8	ICP ICP-MS	100 100	ug/L ug/L
Antimony (Sb)	EPA-200.7 EPA-200.8 EPA-200.9	ICP ICP-MS GFAA		ug/L ug/L ug/L
Arsenic (As)	EPA-200.7 EPA-200.8 EPA-200.9	ICP ICP-MS GFAA		ug/L ug/L ug/L

Table 9.1. Continued...

Parameter	Method Reference	Method Description	Report Limit	ing
			2,,,,,,	
Barium (Ba)	EPA-200.7	ICP	300	ug/L
•	EPA-200.8	ICP-MS	300	ug/l
Beryllium (Be)	EPA-200.7	1CP	5.0	
	EPA-200.8	ICP-MS	1.0	
	EPA-200.9	GFAA	1.0	ug/l
Boron (B)	EPA-200.7	ICP	50	ug/l
Cadmium (Cd)	EPA-200.7	ICP	30.0	ug/l
	EPA-200.8	ICP-MS	1.0	ug/l
	EPA-200.9	GFAA	1.0	ug/L
Calcium (Ca)	EPA-200.7	1CP	1000	ug/L
Chromium (Cr)	EPA-200.7	ICP	40.0	ug/l
	EPA-200.8	ICP-MS		ug/l
	EPA-200.9	GFAA		ug/l
Cobalt (Co)	EPA-200.7	ICP	20.0	ug/l
	EPA-200.8	ICP-MS	5.0	ug/l
Copper (Cu)	EPA-200.7	ICP		ug/l
	EPA-200.8	ICP-MS	50.0	ug/l
ron (Fe)	EPA-200.7	ICP	100	ug/L
lardness	EPA-200.7	Calculation (ICP)	10000	ug/L
ead (Pb)	EPA-200.7	ICP		ug/l
	EPA-200.8	ICP-MS		ug/l
	EPA-200.9	GFAA	5.0	ug/l
lagnesium (Mg)	EPA-200.7	ICP	1000	ug/i
langanese (Mn)	EPA-200.7	ICP	10.0	
	EPA-200.8	ICP-MS	10.0	ug/l
lercury (Hg)	EPA-245.1	Automated Cold Vapor	0.5	
	EPA-200.8	ICP-MS	0.5	ug/l
olybdenum (Mo)	EPA-200.7	ICP	20.0	ug/l
	EPA-200.8	ICP-MS	5.0	
ickel (Ni)	EPA-200.7	ICP	10.0	
	EPA-200.8	ICP-MS	5.0	ug/l
		•		

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• Fable 9.1. Continued...

Silver (Ag)       EPA-200.7 EPA-200.8 ICP-MS 40.0 EPA-200.8 EPA-200.9 GFAA       1CP-MS 40.0 40.0 GFAA         Sodium (Na)       EPA-200.7 ICP 1000         Strontium (Sr)       EPA-200.7 ICP 1000         Thallium (Tl)       EPA-200.8 EPA-200.9 GFAA       1.5 EPA-200.9 GFAA         Tin (Sn)       EPA-200.7 ICP 2000         Vanadium (V)       EPA-200.7 ICP 50.0 ICP-MS 5.0         Zinc (Zn)       EPA-200.7 EPA-200.8 ICP-MS 50.0 ICP-M	Parameter	Method Reference	Method Description	Report Limit	ing
EPA-200.8	· · · · · · · · · · · · · · · · · · ·			· · · · · · · · · · · · · · · · · · ·	
Silver (Ag)       EPA-200.7 EPA-200.8 ICP-MS 40.0 EPA-200.8 EPA-200.9 GFAA       1CP 40.0 EPA-80.0 40.0 GFAA         Sodium (Na)       EPA-200.7 ICP 1000         Strontium (Sr)       EPA-200.7 ICP 1000         Thallium (Tl)       EPA-200.8 EPA-200.9 GFAA       1.5 EPA-200.9 GFAA         Tin (Sn)       EPA-200.7 ICP 2000       2000         Vanadium (V)       EPA-200.7 EPA-200.8 ICP-MS       50.0 EPA-200.8 ICP-MS         Zinc (Zn)       EPA-200.7 EPA-200.8 ICP-MS       50.0 EPA-200.8 ICP-MS         SANICS       EPA-200.8 ICP-MS       50.0 EPA-200.8 ICP-MS         Benzene       EPA 524.2 GC/MS Volatiles       0.5 EPA-200.8 ICP-MS         Bromobenzene       EPA 524.2 GC/MS Volatiles       0.5 EPA-200.8 ICP-MS         Bromodichloromethane       EPA 524.2 GC/MS Volatiles       0.5 EPA-200.8 ICP-MS         Bromoform       EPA 524.2 GC/MS Volatiles       0.5 EPA-200.8 ICP-MS         Bromoform       EPA 524.2 GC/MS Volatiles       0.5 EPA-200.8 ICP-MS         Bromomethane       EPA 524.2 GC/MS Volatiles       0.5 ICP-MS         Bromometha	elenium (Se)	EPA-200.7	ICP	100	ug/l
EPA-200.7   ICP		EPA-200.8	ICP-MS	5.0	ug/
EPA-200.8 EPA-200.9 GFAA 40.0 GFAA 40.0 Sodium (Na) EPA-200.7 ICP 1000  Strontium (Sr) EPA-200.7 ICP 1000  Thallium (Tl) EPA-200.8 ICP-MS 1.5 EPA-200.9 GFAA 1.5  Tin (Sn) EPA-200.7 ICP 2000  Vanadium (V) EPA-200.7 ICP 50.0 ICP-MS 5.0  Zinc (Zn) EPA-200.8 ICP-MS 5.0  ANICS  Benzene EPA 524.2 GC/MS Volatiles 0.5  Bromobenzene EPA 524.2 GC/MS Volatiles 0.5  Bromoform EPA 524.2 GC/MS Volatiles 0.5  Bromoform EPA 524.2 GC/MS Volatiles 0.5  Bromoform EPA 524.2 GC/MS Volatiles 0.5  Bromomethane EPA 524.2 GC/MS Volatiles 0.5		EPA-200.9	GFAA	5.0	ug/
EPA-200.8 EPA-200.9 GFAA 40.0 GFAA 4	ilver (Ag)	EPA-200.7	ICP	40.0	ug/
EPA-200.9       GFAA       40.0         Sodium (Na)       EPA-200.7       ICP       1000         Strontium (Sr)       EPA-200.7       ICP       1000         Thallium (Tl)       EPA-200.8       ICP-MS       1.5         EPA-200.9       GFAA       1.5         Tin (Sn)       EPA-200.7       ICP       2000         Vanadium (V)       EPA-200.7       ICP-MS       50.0         Zinc (2n)       EPA-200.8       ICP-MS       50.0         ANICS       ICP-MS       50.0         Benzene       EPA-200.8       ICP-MS       50.0         Bromochloromethane       EPA 524.2       GC/MS Volatiles       0.5         Bromochloromethane       EPA 524.2       GC/MS Volatiles       0.5         Bromoform       EPA 524.2       GC/MS Volatiles       0.5         Bromomethane       EPA 524.2       GC/MS Volatiles       0.5         Bromomethane       EPA 524.2       GC/MS Volatiles       0.5         tert-Butylbenzene       EPA 524.2       GC/MS Volatiles       0.5         tert-Butylbenzene       EPA 524.2       GC/MS Volatiles       0.5			ICP-MS	40.0	_
Strontium (Sr)		EPA-200.9	GFAA	40.0	
Thallium (TI)	odium (Na)	EPA-200.7	ICP	1000	ug/i
EPA-200.9 GFAA 1.5  Tin (Sn) EPA-200.7 ICP 2000  Vanadium (V) EPA-200.7 ICP 50.0 EPA-200.8 ICP-MS 5.0  Zinc (Zn) EPA-200.8 ICP-MS 50.0 EPA-200.8 ICP-MS 50.0  Benzene EPA 524.2 GC/MS Volatiles 0.5  Bromochloromethane EPA 524.2 GC/MS Volatiles 0.5  Bromodichloromethane EPA 524.2 GC/MS Volatiles 0.5  Bromoform EPA 524.2 GC/MS Volatiles 0.5  Bromomoform EPA 524.2 GC/MS Volatiles 0.5  Bromomomoform EPA 524.2 GC/MS Volatiles 0.5  Bromomomothane EPA 524.2 GC/MS Volatiles 0.5  Bromomethane EPA 524.2 GC/MS Volatiles 0.5  Bromomethane EPA 524.2 GC/MS Volatiles 0.5  Bromomethane EPA 524.2 GC/MS Volatiles 0.5	trontium (Sr)	EPA-200.7	ICP	1000	ug/
EPA-200.9 GFAA 1.5  Tin (Sn) EPA-200.7 ICP 2000  Vanadium (V) EPA-200.7 ICP 50.0	hallium (Tl)	EPA-200.8	ICP-MS	1.5	ug/
Vanadium (V)         EPA-200.7 EPA-200.8 ICP-MS         50.0 EPA-200.8 ICP-MS <td< td=""><td></td><td>EPA-200.9</td><td>GFAA</td><td></td><td>ug/</td></td<>		EPA-200.9	GFAA		ug/
EPA-200.8 ICP-MS 5.0  Zinc (Zn) EPA-200.7 ICP 50.0 EPA-200.8 ICP-MS 50.0  ANICS  Benzene EPA 524.2 GC/MS Volatiles 0.5  Bromobenzene EPA 524.2 GC/MS Volatiles 0.5  Bromochloromethane EPA 524.2 GC/MS Volatiles 0.5  Bromodichloromethane EPA 524.2 GC/MS Volatiles 0.5  Bromoform EPA 524.2 GC/MS Volatiles 0.5  Bromomethane EPA 524.2 GC/MS Volatiles 0.5	in (Sn)	EPA-200.7	ICP	2000	ug/
Zinc (Zn) EPA-200.7 EPA-200.8 ICP 50.0 EPA-200.8 ICP-MS 50.0 ANICS  Benzene EPA 524.2 GC/MS Volatiles 0.5 Bromochloromethane EPA 524.2 GC/MS Volatiles 0.5 Bromodichloromethane EPA 524.2 GC/MS Volatiles 0.5 Bromodichloromethane EPA 524.2 GC/MS Volatiles 0.5 Bromoform EPA 524.2 GC/MS Volatiles 0.5 Bromomethane EPA 524.2 GC/MS Volatiles 0.5 Bromomethane EPA 524.2 GC/MS Volatiles 0.5 Bromomethane EPA 524.2 GC/MS Volatiles 0.5	anadium (V)	EPA-200.7	ICP	50.0	ug/
ANICS  Benzene EPA 524.2 GC/MS Volatiles 0.5  Bromobenzene EPA 524.2 GC/MS Volatiles 0.5  Bromochloromethane EPA 524.2 GC/MS Volatiles 0.5  Bromodichloromethane EPA 524.2 GC/MS Volatiles 0.5  Bromodichloromethane EPA 524.2 GC/MS Volatiles 0.5  Bromoform EPA 524.2 GC/MS Volatiles 0.5  Bromomethane EPA 524.2 GC/MS Volatiles 0.5  Bromomethane EPA 524.2 GC/MS Volatiles 0.5  n-Butylbenzene EPA 524.2 GC/MS Volatiles 0.5  tert-Butylbenzene EPA 524.2 GC/MS Volatiles 0.5		EPA-200.8	ICP-MS	5.0	ug/
Benzene EPA 524.2 GC/MS Volatiles 0.5 Bromobenzene EPA 524.2 GC/MS Volatiles 0.5 Bromochloromethane EPA 524.2 GC/MS Volatiles 0.5 Bromodichloromethane EPA 524.2 GC/MS Volatiles 0.5 Bromoform EPA 524.2 GC/MS Volatiles 0.5 Bromomethane EPA 524.2 GC/MS Volatiles 0.5	inc (Zn)	EPA-200.7	ICP	50.0	ug/
Benzene EPA 524.2 GC/MS Volatiles 0.5 Bromobenzene EPA 524.2 GC/MS Volatiles 0.5 Bromochloromethane EPA 524.2 GC/MS Volatiles 0.5 Bromodichloromethane EPA 524.2 GC/MS Volatiles 0.5 Bromoform EPA 524.2 GC/MS Volatiles 0.5 Bromomethane EPA 524.2 GC/MS Volatiles 0.5 Bromomethane EPA 524.2 GC/MS Volatiles 0.5 Bromomethane EPA 524.2 GC/MS Volatiles 0.5 br-Butylbenzene EPA 524.2 GC/MS Volatiles 0.5 tert-Butylbenzene EPA 524.2 GC/MS Volatiles 0.5	NICC	EPA-200.8	ICP-MS	50.0	ug/
Bromobenzene EPA 524.2 GC/MS Volatiles 0.5 Bromochloromethane EPA 524.2 GC/MS Volatiles 0.5 Bromodichloromethane EPA 524.2 GC/MS Volatiles 0.5 Bromoform EPA 524.2 GC/MS Volatiles 0.5 Bromomethane EPA 524.2 GC/MS Volatiles 0.5	n1C3				
Bromochloromethane EPA 524.2 GC/MS Volatiles 0.5 Bromodichloromethane EPA 524.2 GC/MS Volatiles 0.5 Bromoform EPA 524.2 GC/MS Volatiles 0.5 Bromomethane EPA 524.2 GC/MS Volatiles 0.5 n-Butylbenzene EPA 524.2 GC/MS Volatiles 0.5 tert-Butylbenzene EPA 524.2 GC/MS Volatiles 0.5	enzene	EPA 524.2	GC/MS Volatiles	0.5	ug/
Bromodichloromethane EPA 524.2 GC/MS Volatiles 0.5 Bromoform EPA 524.2 GC/MS Volatiles 0.5 Bromomethane EPA 524.2 GC/MS Volatiles 0.5 n-Butylbenzene EPA 524.2 GC/MS Volatiles 0.5 tert-Butylbenzene EPA 524.2 GC/MS Volatiles 0.5	romobenzene	EPA 524.2	GC/MS Volatiles	0.5	ug/
Bromoform EPA 524.2 GC/MS Volatiles 0.5 Bromomethane EPA 524.2 GC/MS Volatiles 0.5 n-Butylbenzene EPA 524.2 GC/MS Volatiles 0.5 tert-Butylbenzene EPA 524.2 GC/MS Volatiles 0.5	romochloromethane	EPA 524.2	GC/MS Volatiles	0.5	ug/
Bromomethane EPA 524.2 GC/MS Volatiles 0.5 n-Butylbenzene EPA 524.2 GC/MS Volatiles 0.5 tert-Butylbenzene EPA 524.2 GC/MS Volatiles 0.5	romodichloromethane	EPA 524.2	GC/MS Volatiles	0.5	ug/i
n-Butylbenzene EPA 524.2 GC/MS Volatiles 0.5 tert-Butylbenzene EPA 524.2 GC/MS Volatiles 0.5	romoform	EPA 524.2	GC/MS Volatiles	0.5	ug/
tert-Butylbenzene EPA 524.2 GC/MS Volatiles 0.5	romomethane	EPA 524.2	GC/MS Volatiles	0.5	ug/
	-Butylbenzene	EPA 524.2	GC/MS Volatiles	0.5	ug/
sec-Butylbenzene EPA 524.2 GC/MS Volatiles 0.5	ert-Butylbenzene	EPA 524.2	GC/MS Volatiles	0.5	ug/
	ec-Butylbenzene	EPA 524.2	GC/MS Volatiles	0.5	ug/
Carbon Tetrachloride EPA 524.2 GC/MS Volatiles 0.5	amban Tabanahla-ida	EDA 52/ 2	CC/Me Valetilas	0.5	

Table 9.1. Continued...

	<u> </u>		
Parameter	Method Reference	Method Description	Reporting Limit
Chlorobenzene	EPA 524.2	GC/MS Volatiles	0.5 ug/L
Chloroethane	EPA 524.2	GC/MS Volatiles	0.5 ug/L
Chloroform	EPA 524.2	GC/MS Volatiles	0.5 ug/L
Chloromethane	EPA 524.2	GC/MS Volatiles ~	0.5 ug/L
o-Chlorotoluene	EPA 524.2	GC/MS Volatiles	0.5 ug/L
p-Chlorotoluene	EPA 524.2	GC/MS Volatiles	0.5 ug/L
Dibromochloromethane	EPA 524.2	GC/MS Volatiles	0.5 ug/L
Dibromomethane	EPA 524.2	GC/MS Volatiles	0.5 ug/L
1,2-Dichlorobenzene	EPA 524.2	GC/MS Volatiles	0.5 ug/L
1,3-Dichlorobenzene	EPA 524.2	GC/MS Volatiles	0.5 ug/L
1,4-Dichlorobenzene	EPA 524.2	GC/MS Volatiles	0.5 ug/L
1,1-Dichloroethane	EPA 524.2	GC/MS Volatiles	0.5 ug/L
1,2-Dichloroethane	EPA 524.2	GC/MS Volatiles	0.5 ug/L
1,1-Dichloroethene	EPA 524.2	GC/MS Volatiles	0.5 ug/L
cis-1,2,Dichloroethene	EPA 524.2	GC/MS Volatiles	0.5 ug/L
trans-1,2-Dichloroethene	EPA 524.2	GC/MS Volatiles	0.5 ug/L
1,1-Dichloropropene	EPA 524.2	GC/MS Volatiles	0.5 ug/L
1,2-Dichloropropane	EPA 524.2	GC/MS Volatiles	0.5 ug/L
cis-1,3-Dichloropropene	EPA 524.2	GC/MS Volatiles	0.5 ug/L
trans-1,3-Dichloropropene	EPA 524.2	GC/MS Volatiles	0.5 ug/L
1,3-Dichloropropane	EPA 524.2	GC/MS Volatiles	0.5 ug/L
2,2-Dichloropropane	EPA 524.2	GC/MS Volatiles	0.5 ug/L
Ethyl benzene	EPA 524.2	GC/MS Volatiles	0.5 ug/L
Fluorotrichloromethane	EPA 524.2	GC/MS Volatiles	0.5 ug/L
Hexachlorobutadiene	EPA 524.2	GC/MS Volatiles	0.5 ug/L
Isopropylbenzene	EPA 524.2	GC/MS Volatiles	0.5 ug/L

Table 9.1. Continued...

Parameter	Method Reference	Method Description	Reporting Limit
p-Isopropyltoluene	EPA 524.2	GC/MS Volatiles	0.5 ug/L
Methylene Chloride	EPA 524.2	GC/MS Volatiles	0.5 ug/L
Naphthalene	EPA 524.2	GC/MS Volatiles	0.5 ug/L
n-Propylbenzene	EPA 524.2	GC/MS Volatiles -	0.5 ug/L
Styrene	EPA 524.2	GC/MS Volatiles	0.5 ug/L
1,1,1,2-Tetrachloroethane	EPA 524.2	GC/MS Volatiles	0.5 ug/L
1,1,2,2-Tetrachloroethane	EPA 524.2	GC/MS Volatiles	0.5 ug/L
1,2,3-Trichloropropane	EPA 524.2	GC/MS Volatiles	0.5 ug/L
1,2,4-Trichlorobenzene	EPA 524.2	GC/MS Volatiles	0.5 ug/L
1,2,3-Trichlorobenzene	EPA 524.2	GC/MS Volatiles	0.5 ug/L
1,2,4-Trimethylbenzene	EPA 524.2	GC/MS Volatiles	0.5 ug/L
1,3,5-Trimethylbenzene	EPA 524.2	GC/MS Volatiles	0.5 ug/L
Tetrachloroethene	EPA 524.2	GC/MS Volatiles	0.5 ug/L
Toluene	EPA 524.2	GC/MS Volatiles	0.5 ug/L
1,1,1-Trichloroethane	EPA 524.2	GC/MS Volatiles	0.5 ug/L
1,1,2-Trichloroethane	EPA 524.2	GC/MS Volatiles	0.5 ug/L
Trichloroethene	EPA 524.2	GC/MS Volatiles	0.5 ug/L
/inyl Chloride	EPA 524.2	GC/MS Volatiles	0.5 ug/L
p-Xylene	EPA 524.2	GC/MS Volatiles	0.5 ug/L
n & p Xylene	EPA 524.2	GC/MS Volatiles	0.5 ug/L
Xylenes, total	EPA 524.2	GC/MS Volatiles	0.5 ug/L

del des Table 9.2. Analytical Methods and Reporting Limits - RCRA

Parameter	Method Reference	Method: Description	Reporti	ng Limit
rai ameter	Reference	besch iperon	Aqueous	Non-Aqueous
Vet Chemistry				
Cyanide, Amenable	SW-9012	Spectrophotometric	0.005 mg/L	0.125 mg/Kg
Cyanide, Total	SW-9012	Spectrophotometric	0.005 mg/L	0.125 mg/Kg
Hexavalent Chromium	SW-7196A	Colorimetric	0.010 mg/L	5.0 mg/Kg
Ignitability	sw-1010	Pensky Martins	NA	NA
Oil & Grease	sw-9070	Gravimetric	5.0 mg/L	
Paint Filter Test	SW-9095A	NA	NA	NA
рн .	SW-9040B, SW-9041A, SW-9045C	Potentiometric, pH Paper	NA	NA
TCLP Extraction	sw-1311	18 hr Extraction	NA	NA
fetals				
Aluminum (Al)	SW-6010A SW-6020	ICP ICP-MS	0.10 mg/L 0.050 mg/L	5.0 mg/Kg 50.0 mg/Kg
Antimony (Sb)	SW-6010A	ICP	0.10 mg/L	5.0 mg/Kg
, , , , , , , , , , , , , , , , , , ,	SW-6020 SW-7041	ICP-MS GFAA	0.001 mg/L 0.020 mg/L	1.0 mg/Kg 1.0 mg/Kg
Arsenic (As)	SW-6010A	ICP	0.10 mg/L	5.0 mg/Kg
A COLLEGE (AG)	SW-6020	ICP-MS	0.005 mg/L	5.0 mg/Kg
	SW-7060A	GFAA	0.005 mg/L	0.25 mg/Kg
Barium (Ba)	SW-6010A	ICP	0.020 mg/L	1.0 mg/Kg
	SW-6020	ICP-MS	0.005 mg/L	5.0 mg/Kg
Beryllium (Be)	SW-6010A	ICP	0.005 mg/L	0.25 mg/Kg
	sw-6020	ICP-MS	0.001 mg/L	1.0 mg/Kg
	SW-7091	GFAA	0.001 mg/L	0.05 mg/Kg
Boron (B)	SW-6010A	ICP	0.050 mg/L	2.5 mg/Kg
Cadmium (Cd)	SW-6010A	ICP	0.030 mg/L	1.5 mg/Kg
	sw-6020	ICP-MS	0.001 mg/L	1.0 mg/Kg
	SW-7131A	GFAA	0.001 mg/L	0.05 mg/Kg
Calcium (Ca)	SW-6010A	ICP	1.0 mg/L	50.0 mg/Kg
Chromium (Cr)	SW-6010A	ICP	0.040 mg/L	2.0 mg/Kg
	sw-6020	ICP-MS	0.002 mg/L	2.0 mg/Kg
	sw-7191	GFAA	0.002 mg/L	0.1 mg/Kg

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Table 9.2. Continued

Parameter	Method Reference	Method Description	1	Reportir	ng Limit	
		* **	Aque	ous	Non-	-Aqueou
Cobalt (Co)	SW-6010A	ICP	0.020	mg/L		mg/Kg
·•	sw-6020	ICP-MS	0.005	mg/L	5.0	mg/Kg
	sw-7201	GFAA	0.005	mg/L	0.25	mg/Kg
Copper (Cu)	SW-6010A	ICP	0.020	mg/L	1.0	mg/Kg
	sw-6020	1CP-MS	0-005	mg/L	5.0	mg/Kg
Iron (Fe)	SW-6010A	ICP	0.10	mg/L	5.0	mg/Kg
Lead (Pb)	SW-6010A	ICP	0.080	mg/L	4.0	mg/Kg
	sw-6020	ICP-MS	0.001	mg/L	1.0	mg/Kg
	sw-7421	GFAA	0.005			mg/Kg
Magnesium (Mg)	SW-6010A	ICP	1.0	mg/L	50.0	mg/Kg
Manganese (Mn)	SW-6010A	ICP	0.010	mg/L	0.50	mg/Kg
	sw-6020	ICP-MS	0.010			mg/Kg
Mercury (Hg)	SW-7470A/SW-7471A	Automated Cold Vapor	0.0002	mg/L	0.01	mg/Kg
Molybdenum (Mo)	SW-6010A	ICP	0.020	mg/L	1.0	mg/Kg
	sw-6020	ICP-MS	0.001	mg/L	1.0	mg/Kg
Nickel (Ni)	SW-6010A	ICP	0.010		0.5	mg/Kg
	sw-6020	ICP-MS	0.005	mg/L	5.0	mg/Kg
Potassium (K)	SW-6010A	ICP	1.0	mg/L	50.0	mg/Kg
Selenium (Se)	SW-6010A	ICP	0.10	mg/L	5.0	mg/Kg
	sw-6020	ICP-MS	0.005	mg/L	5.0	mg/Kg
	sw-7740	GFAA	0.005	mg/L	0.25	mg/Kg
Silver (Ag)	SW-6010A	ICP	0.040	mg/L	2.0	mg/Kg
	sw-6020	ICP-MS	0.0005	mg/L	0.5	mg/Kg
	sw-7761	GFAA	0.001	mg/L	0.05	mg/Kg
Sodium (Na)	SW-6010A	ICP	1.0	mg/L	50.0	mg/Kg
Strontium (Sr)	SW-6010A	1CP	0.10	mg/L	5.0	mg/Kg
Thallium (Tl)	SW-6010A	ICP	0.50	mg/L	25	mg/Kg
	sw-6020	ICP-MS	0.001		1.0	mg/Kg
	sw-7841	GFAA	0.010	mg/L	0.5	mg/Kg
Tin (Sn)	SW-6010A	1CP	2.0	mg/L	100	mg/Kg
	SW-6010A	ICP		mg/L		mg/Kg

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Table 9.2. Continued

Parameter	Method Reference	Method Description	R	leporting	Limit	
r at affecter	Reference	besch iperon	Aqueo	ous	Non	-Aqueous
Vanadium (V)	SW-6010A SW-6020	ICP ICP-MS	0.050 0.005	<del>-</del> -		mg/Kg mg/Kg
Zinc (Zn)	SW-6010A SW-6020	ICP ICP-MS	0.050 0.050			mg/Kg mg/Kg
Organics - Volatile Compounds						
Acetone	SW-8260A	GC/MS	20	ug/L	100	ug/Kg
Acrolein	SW-8260A	GC/MS	50	ug/L	50	ug/Kg
Acrylonitrile	SW-8260A	GC/MS	50	ug/L	50	ug/Kg
Allyl chloride	SW-8260A	GC/MS	5.0	ug/L	5.0	ug/Kg
Benzene	SW-8260A	GC/MS	1.0	ug/L	5.0	ug/Kg
Bromobenzene	SW-8260A	GC/MS	1.0	ug/L	5.0	ug/Kg
Bromochloromethane	SW-8260A	GC/MS	1.0	ug/L	5.0	ug/Kg
Bromodichloromethane	SW-8260A	GC/MS	1.0	ug/L	5.0	ug/Kg
Bromoform	SW-8260A	GC/MS	1.0	ug/L	5.0	ug/Kg
Bromomethane	SW-8260A	GC/MS	5.0	ug/L	5.0	ug/Kg
n-Butylbenzene	SW-8260A	GC/MS	1.0	ug/L	5.0	ug/Kg
tert-Butylbenzene	SW-8260A	GC/MS	1.0	ug/L	5.0	ug/Kg
sec-Butylbenzene	SW-8260A	GC/MS	1.0	ug/L	5.0	ug/Kg
2-Butanone (MEK)	SW-8260A	GC/MS	20	ug/L	100	ug/Kg
Carbon Disulfide	SW-8260A	GC/MS	1.0	ug/L	5.0	ug/Kg
Carbon Tetrachloride	SW-8260A	GC/MS	1.0	ug/L	5.0	ug/Kg
Chlorobenzene	SW-8260A	GC/MS	1.0	ug/L	5.0	ug/Kg
Chloroethane	SW-8260A	GC/MS	1.0	ug/L	5.0	ug/Kg
2-Chloroethylvinyl ether	SW-8260A	GC/MS	5.0	ug/L	5.0	ug/Kg
Chloroform	SW-8260A	GC/MS	1.0	ug/L	5.0	ug/Kg
Chloromethane	SW-8260A	GC/MS	5.0	ug/L	10.0	ug/Kg

Danamatan	Method	Method		Reportin	g Limit	
Parameter	Reference	Description	Aque	eous	Non	-Aqueous
Chloroprene	SW-8260A	GC/MS	5.0	ug/L	5.0	ug/Kg
o-Chlorotoluene	SW-8260A	GC/MS	1.0	ug/L	5.0	ug/Kg
p-Chlorotoluene	SW-8260A	GC/MS	1.0	ug/L	5.0	ug/Kg
Dibromochloromethane	SW-8260A	GC/MS	1.0	ug/L	5.0	ug/Kg
1,2-Dibromo-3-Chloropropane	SW-8260A	GC/MS	5.0	ug/L	5.0	ug/Kg
1,2-Dibromoethane	SW-8260A	GC/MS	5.0	ug/L	5.0	ug/Kg
Dibromomethane	SW-8260A	GC/MS	5.0	ug/L	5.0	ug/Kg
1,2-Dichlorobenzene	SW-8260A	GC/MS	1.0	ug/L	5.0	ug/Kg
1,3-Dichlorobenzene	SW-8260A	GC/MS	1.0	ug/L	5.0	ug/Kg
1,4-Dichlorobenzene	SW-8260A	GC/MS	1.0	ug/L	5.0	ug/Kg
trans-1,4-Dichloro-2-butene	SW-8260A	GC/MS	5.0	ug/L	5.0	ug/Kg
Dichlorodifluoromethane	SW-8260A	GC/MS	1.0	ug/L	5.0	ug/Kg
1,1-Dichloroethane	SW-8260A	GC/MS	1.0	ug/L	5.0	ug/Kg
1,2-Dichloroethane	SW-8260A	GC/MS	1.0	ug/L	5.0	ug/Kg
1,1-Dichloroethene	SW-8260A	GC/MS	1.0	ug/L	5.0	ug/Kg
cis-1,2,Dichloroethene	SW-8260A	GC/MS	1.0	ug/L	5.0	ug/Kg
trans-1,2-Dichloroethene	SW-8260A	GC/MS	1.0	ug/L	5.0	ug/Kg
1,1-Dichloropropene	SW-8260A	GC/MS	1.0	ug/L	5.0	ug/Kg
1,2-Dichloropropane	SW-8260A	GC/MS	1.0	ug/L	5.0	ug/Kg
cis-1,3-Dichloropropene	SW-8260A	GC/MS	1.0	ug/L	5.0	ug/Kg
trans-1,3-Dichloropropene	SW-8260A	GC/MS	1.0	ug/L	5.0	ug/Kg
1,3-Dichloropropane	SW-8260A	GC/MS	1.0	ug/L	5.0	ug/Kg
2,2-Dichloropropane	SW-8260A	GC/MS	1.0	ug/L	5.0	ug/Kg
Ethyl benzene	SW-8260A	. GC/MS	1.0	ug/L	5.0	ug/Kg
Ethyl methacrylate	SW-8260A	GC/MS	1.0	ug/L	5.0	ug/Kg

Parameter	Method Method	Method Method Report Reference Description		Reportir	ng Limit	
r di difecter	Ke fer ende	2	upA	eous	Non	-Aqueous
Fluorotrichlöromethane	SW-8260A	GC/MS	1.0	ug/L	5.0	ug/Kg
Hexach lorobutadiene	SW-8260A	GC/MS	5.0	ug/L	5.0	ug/Kg
2-Hexanone	SW-8260A	GC/MS	<sub>~</sub> 10	ug/L	50	ug/Kg
Iodomethane	SW-8260A	GC/MS	5.0	ug/L	5.0	ug/Kg
Isopropylbenzene	SW-8260A	GC/MS	1.0	ug/L	5.0	ug/Kg
p-Isopropyltoluene	SW-8260A	GC/MS	1.0	ug/L	5.0	ug/Kg
Methacrylonitrile	SW-8260A	GC/MS	1.0	ug/L	5.0	ug/Kg
Methylene Chloride	SW-8260A	GC/MS	10	ug/L	10	ug/Kg
4-Methyl-2-pentanone (MIBK)	SW-8260A	GC/MS	10	ug/L	50	ug/Kg
Methyl tert butyl ether (MTBE)	SW-8260A	GC/MS	1.0	ug/L	5.0	ug/Kg
Methyl methacrylate	SW-8260A	GC/MS	1.0	ug/L	5.0	ug/Kg
laphthalene	SW-8260A	GC/MS	5.0	ug/L	5.0	ug/Kg
Propionitrile	SW-8260A	GC/MS	50	ug/L	50	ug/Kg
n-Propylbenzene	SW-8260A	GC/MS	1.0	ug/L	5.0	ug/Kg
Styrene	SW-8260A	GC/MS	1.0	ug/L	5.0	ug/Kg
1,1,1,2-Tetrachloroethane	SW-8260A	GC/MS	1.0	ug/L	5.0	ug/Kg
,1,2,2-Tetrachloroethane	SW-8260A	GC/MS	1.0	ug/L	5.0	ug/Kg
1,2,3-Trichloropropane	SW-8260A	GC/MS	5.0	ug/L	5.0	ug/Kg
1,2,4-Trichlorobenzene	SW-8260A	GC/MS	5.0	ug/L	5.0	ug/Kg
1,2,3-Trichlorobenzene	SW-8260A	GC/MS	5.0	ug/L	5.0	ug/Kg
1,2,4-Trimethylbenzene	SW-8260A	GC/MS	1.0	ug/L	5.0	ug/Kg
1,3,5-Trimethylbenzene	SW-8260A	GC/MS	1.0	ug/L	5.0	ug/Kg
etrachloroethene	SW-8260A	GC/MS	1.0	ug/L	5.0	ug/Kg
oluene	SW-8260A	GC/MS	. 1.0	ug/L	5.0	ug/Kg
I,1,1-Trichloroethane	SW-8260A	GC/MS	1.0	ug/L	5.0	ug/Kg

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Table 9.2. Continued

Parameter	Method Reference	Method : Description		Report	ing Limit	:
rajametei	Reference	besch iperon	Aqu	eous	Nor	-Aqueous
1,1,2-Trichloroethane	SW-8260A	GC/MS	1.0	ug/L	5.0	ug/Kg
Trichloroethene	SW-8260A	GC/MS	1.0	ug/L	5.0	ug/Kg
Vinyl Acetate	SW-8260A	GC/MS	5.0	ug/L	5.0	ug/Kg
Vinyl Chloride	SW-8260A	GC/MS	2.0	ug/L	2.0	ug/Kg
o-Xylene	SW-8260A	GC/MS	1.0	ug/L	5.0	ug/Kg
m & p Xylene	SW-8260A	GC/MS	1.0	ug/L	5.0	ug/Kg
Xylenes, total	SW-8260A	GC/MS	1.0	ug/L	5.0	ug/Kg
n-Hexane	SW-8260A	GC/MS	10	ug/L	10	ug/Kg
EMI-VOLATILE COMPOUNDS						
Acenaphthene	sw-8270B	GC/MS	10	ug/L	330	ug/Kg
Acenaphthylene	SW-8270B	GC/MS	10	ug/L	330	ug/Kg
Acetophenone	SW-8270B	GC/MS	20	ug/L	660	ug/Kg
2-Acetylaminoflourene (2-AAF)	SW-8270B	GC/MS	20	ug/L	660	ug/Kg
4-Aminobipheyl	sw-8270B	GC/MS	20	ug/L	660	ug/Kg
Aniline	SW-8270B	GC/MS	10	ug/L	330	ug/Kg
Anthracene	SW-8270B	GC/MS	10	ug/L	330	ug/Kg
Aramite	SW-8270B	GC/MS	15	ug/L	495	ug/Kg
Benzidine	sw-8270B	GC/MS	50	ug/L	1,650	ug/Kg
Benzo(a)anthracene	SW-8270B	GC/MS	10	ug/L	330	ug/Kg
Benzo(b)fluoranthene	SW-8270B	GC/WS	10	ug/L	330	ug/Kg
Benzo(k)fluoranthene	SW-8270B	GC/MS	10	ug/L	330	ug/Kg
Benzo(a)pyrene	SW-8270B	GC/MS	10	ug/L	330	ug/Kg
Benzo(g,h,i)perylene	SW-8270B	GC/MS	10	ug/L	330	ug/Kg
Benzyl alcohol	SW-8270B	GC/MS	10	ug/L	330	ug/Kg
Benzyl butyl phthalate	SW-8270B	GC/MS	10	ug/L	330	ug/Kg

Parameter	Method Reference	Method: Description		Report	ing Limi	t
r di ameter	Reference		Aqu	ieous	No.	n-Aqueous
Bis(2-chloroethyl)ether	SW-8270B	GC/MS	10	ug/L	330	ug/Kg
Bis(2-chloroethoxy)methane	SW-8270B	GC/MS	10	ug/L	330	ug/Kg
Bis(2-ethylhexyl)phthalate	SW-8270B	GC/MS	40	ug/L	330	ug/Kg
Bis(2-chloroisopropyl)ether	SW-8270B	GC/MS	10	ug/L	330	ug/Kg
4-Bromophenyl phenyl ether	SW-8270B	GC/MS	10	ug/L	330	ug/Kg
4-Chl <i>o</i> roaniline	SW-8270B	GC/MS	10	ug/L	330	ug/Kg
Chlorobenzilate	SW-8270B	GC/MS	50	ug/L	1,650	ug/Kg
2-Chloronaphthalene	SW-8270B	GC/MS	10	ug/L	330	ug/Kg
4-Chlorophenyl phenyl ether	SW-8270B	GC/MS	10	ug/L	330	ug/Kg
Chrysene	SW-8270B	GC/MS	10	ug/L	330	ug/Kg
Diallate	SW-8270B	GC/MS	30	ug/L	990	ug/Kg
Dibenzo(a,h)anthracene	SW-8270B	GC/MS	10	ug/L	330	ug/Kg
Dibenzofuran	SW-8270B	GC/MS	10	ug/L	330	ug/Kg
Di-n-butylphthalate	SW-8270B	GC/MS	10	ug/L	330	ug/Kg
1,2-Dichlorobenzene	SW-8270B	GC/MS	10	ug/L	330	ug/Kg
1,3-Dichlorobenzene	SW-8270B	GC/MS	10	ug/L	330	ug/Kg
1,4-Dichlorobenzene	SW-8270B	GC/MS	10	ug/L	330	ug/Kg
3,3-Dichlorobenzidine	SW-8270B	GC/MS	50	ug/L	1,650	ug/Kg
Diethyl phthalate	SW-8270B	GC/MS	10	ug/L	330	ug/Kg
Dimethoate	SW-8270B	GC/MS	100	ug/L	3,300	ug/Kg
o-(Dimethylamino)-azobenzene	SW-8270B	GC/MS	50	ug/L	1,650	ug/Kg
7,12-Dimethylbenz(a)anthracene	SW-8270B	GC/MS	20	ug/L	660	ug/Kg
3,3'-Dimethylbenzidine	SW-8270B	GC/MS	20	ug/L	660	ug/Kg
a,a-Dimethyl-phenethylamino	SW-8270B	GC/MS	50	ug/L	1,650	ug/Kg
Dimethyl phthalate	SW-8270B	GC/MS	10	ug/L	330	ug/Kg

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Table 9.2. Continued

Parameter	Method Reference	Method: Description		Report	ing Limi	t
			Aqu	Jeous	Noi	n-Aqueous
2,4-Dinitrotõluene	SW-8270B	GC/MS	10	ug/L	330	ug/Kg
2,6-Dinitrotoluene	SW-8270B	GC/MS	10	ug/L	330	ug/Kg
Di-n-octylphthalate	sw-8270B	GC/MS	40	ug/L	330	ug/Kg
Diphenylhydrazine	SW-8270B	GC/MS	10	ug/L	330	ug/Kg
Diphenylamine	SW-8270B	GC/MS	20	ug/L	660	ug/Kg
Disulfoton	SW-8270B	GC/MS	20	ug/L	660	ug/Kg
Ethyl methanesulfonate	SW-8270B	GC/MS	20	ug/L	660	ug/Kg
Famphur	SW-8270B	GC/MS	10	ug/L	330	ug/Kg
Fluoranthene	SW-8270B	GC/MS	10	ug/L	330	ug/Kg
Fluorene	SW-8270B	GC/MS	10	ug/L	330	ug/Kg
Hexachlorobenzene	SW-8270B	GC/MS	10	ug/L	330	ug/Kg
Hexachlorobutadiene	SW-8270B	GC/MS	10	ug/L	330	ug/Kg
Hexachlorocyclopentadiene	SW-8270B	GC/MS	10	ug/L	330	ug/Kg
Hexachloroethane	SW-8270B	GC/MS	10	ug/L	330	ug/Kg
Hexach lorophene	SW-8270B	GC/MS	500	ug/L	16,500	ug/Kg
Hexach loropropene	SW-8270B	GC/MS	30	ug/L	990	ug/Kg
Indeno(1,2,3-cd)pyrene	SW-8270B	GC/MS	10	ug/L	330	ug/Kg
Isodrin	SW-8270B	GC/MS	30	ug/L	990	ug/Kg
Isophorone	SW-8270B	GC/MS	10	ug/L	330	ug/Kg
Isosafrole	SW-82708	GC/MS	20	ug/L	660	ug/Kg
Kepone	SW-8270B	GC/MS	250	ug/L	8,250	ug/Kg
Methapryilene	SW-8270B	GC/MS	100	ug/L	3,300	ug/Kg
3-Methylcholanthrene	SW-8270B	GC/MS	30	ug/L	990	ug/Kg
Methyl methanesulfonate	SW-8270B	GC/MS	20	ug/L	660	ug/Kg
2-Methylnapthalene	SW-8270B	GC/MS	10	ug/L	330	ug/Kg

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Table 9.2. Continued

Parameter	Method Reference	Method: Description		Report	ing Limi	t
rai ametei	Kerel elice	·	Aqu	ieous	No	n-Aqueous
Methyl parathion	sw-8270B	GC/MS	20	ug/L	660	ug/Kg
Naphthalene	SW-8270B	GC/MS	10	ug/L	330	ug/Kg
1,4 Napthoquinone	SW-8270B	GC/MS	100	ug/L	3,300	ug/Kg
1-Napthylamine	SW-8270B	GC/MS	30	ug/L	990	ug/Kg
2-Napthylamine	SW-8270B	GC/MS	30	ug/L	990	ug/Kg
Nitrobenzene	sw-8270B	GC/MS	10	ug/L	330	ug/Kg
2-Nitroaniline	SW-8270B	GC/MS	15	ug/L	495	ug/Kg
3-Nitroaniline	SW-8270B	GC/MS	15	ug/L	495	ug/Kg
4-Nitroaniline	SW-8270B	GC/MS	15	ug/L	495	ug/Kg
4-Nitroquinoline-1-oxide	SW-8270B	GC/MS	30	ug/L	990	ug/Kg
N-Nitrosodi-n-butylamine	SW-8270B	GC/MS	20	ug/L	660	ug/Kg
N-Nitrosodiethylamine	SW-8270B	GC/MS	30	ug/L	660	ug/Kg
N-Nitrosodimethylamine	SW-8270B	GC/MS	10	ug/L	330	ug/Kg
N-Nitrosodiphenylamine	sw-8270B	GC/MS	10	ug/L	330	ug/Kg
N-Nitrosodipropylamine	SW-8270B	GC/MS	10	ug/L	330	ug/Kg
N-Nitrosomethylethylamine	sw-8270B	GC/MS	20	ug/L	660	ug/Kg
N-Nitrosomorpholine	SW-8270B	GC/MS	20	ug/L	660	ug/Kg
N-Nitrosopiperidine	sw-8270B	GC/MS	20	ug/L	660	ug/Kg
N-Nitrosopyrrolidine	sw-8270B	GC/MS	20	ug/L	660	ug/Kg
5-Nitro-o-toluidine	sw-8270B	GC/MS	20	ug/L	660	ug/Kg
Parathion	sw-8270B	GC/MS	20	ug/L	660	ug/Kg
Pentachlorobenzene	SW-8270B	GC/MS	20	ug/L	660	ug/Kg
Pentachloronitrobenzene	sw-8270B	GC/MS	20	ug/L	660	ug/Kg
Phenacetin	sw-8270B	GC/MS	20	ug/L	660	ug/Kg
Phenanthrene	SW-8270B	GC/MS	10	ug/L	330	ug/Kg

December	Method	Method:		Report	ing Limit	t
Parameter	Reference	Description 	Aqu	eous	Nor	n-Aqueous
p-Phenylenedïamine	sw-8270B	GC/MS	30	ug/L	990	ug/Kg
Phorate	sw-8270B	GC/MS	20	ug/L	660	ug/Kg
2-Picoline	SW-8270B	GC/MS	20	ug/L	660	ug/Kg
Pronamide	SW-8270B	GC/MS	20	ug/L	660	ug/Kg
Pyrene	SW-8270B	GC/MS	10	ug/L	330	ug/Kg
Pyridine	SW-8270B	GC/MS	10	ug/L	330	ug/Kg
Safrole	SW-8270B	GC/MS	20	ug/L	660	ug/Kg
Sulfotepp	SW-8270B	GC/MS	20	ug/L	660	ug/Kg
1,2,4,5-Tetrachlorobenzene	SW-8270B	GC/MS	20	ug/L	660	ug/Kg
Thionazin	SW-8270B	GC/MS	20	ug/L	660	ug/Kg
o-Toluidine	SW-8270B	GC/MS	20	ug/L	660	ug/Kg
1,2,4-Trichlorobenzene	SW-8270B	GC/MS	10	ug/L	330	ug/Kg
Triethyl phosphorothioate	SW-8270B	GC/MS	20	ug/L	660	ug/Kg
1,3,5-Trinitrobenzene	SW-8270B	GC/MS	30	ug/L	990	ug/Kg
Benzoic Acid	SW-8270B	GC/MS	50	ug/L	1,650	ug/Kg
4-Chloro-3-methylphenol	SW-8270B	GC/MS	10	ug/L	330	ug/Kg
2-Chlorophenol	SW-8270B	GC/MS	10	ug/L	330	ug/Kg
2,4-Dichlorophenol	SW-8270B	GC/MS	10	ug/L	330	ug/Kg
2,6-Dichlorophenol	SW-8270B	GC/MS	10	ug/L	330	ug/Kg
2,4-Dimethylphenol	SW-8270B	GC/MS	10	ug/L	330	ug/Kg
2,4-Dinitrophenol	SW-8270B	GC/MS	10	ug/L	330	ug/Kg
2-Methyl-4,6-dinitrophenol	SW-8270B	GC/MS	10	ug/L	330	ug/Kg
2-Nitrophenol	SW-8270B	GC/MS	10	ug/L	330	ug/Kg
4-Nitrophenol	SW-8270B	GC/MS	10	ug/L	330	ug/Kg
Pentachlorophenol	SW-8270B	GC/MS	10	ug/L	330	ug/Kg

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Table 9.2. Continued

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Parameter	Method Reference	Method Description		Reportir		
		· · · · · · · · · · · · · · · · · · ·	Aque	eous	Nor	n-Aqueous
Phenol :	sw-8270B	GC/MS	10	ug/L	330	ug/Kg
2,3,4,6-Tetrachlorophenol	SW-8270B	GC/MS	20	ug/L	660	ug/Kg
2,4,5-Trichlorophenol	\$W-8270B	GC/MS	-10	ug/L	330	ug/Kg
2,4,6-Trichlorophenol	SW-8270B	GC/MS	10	ug/L	330	ug/Kg
2-Methylphenol	SW-8270B	GC/MS	10	ug/L	330	ug/Kg
3 & 4-Methylphenol	SW-8270B	GC/MS	. 10	ug/L	330	ug/Kg
Pesticides/PCBs						
Aldrin	SH-8080A	GC/ECD	0.2	ug/L	500	ug/Kg
Chlordane	SW-8080A	GC/ECD	0.2	ug/L	500	ug/Kg
Dieldrin	SW-8080A	GC/ECD	0.2	ug/L	500	ug/Kg
4,4'-DDD	SW-8080A	GC/ECD	0.2	ug/L	500	ug/Kg
4,4'-DDE	SW-8080A	GC/ECD	0.2	ug/L	500	ug/Kg
4,4'-DDT	SW-8080A	GC/ECD	0.2	ug/L	500	ug/Kg
Endosulfan I	SW-8080A	GC/ECD	0.2	ug/L	500	ug/Kg
Endosulfan II	SW-8080A	GC/ECD	0.2	ug/L	500	ug/Kg
Endosulfan Sulfate	SW-8080A	GC/ECD	0.2	ug/L	500	ug/Kg
Endrin	SW-8080A	GC/ECD	0.2	ug/L	500	ug/Kg
Endrin Aldehyde	SW-8080A	GC/ECD	0.2	ug/L	500	ug/Kg
Endrin Ketone	A0808-W2	GC/ECD	0.2	ug/L	500	ug/Kg
Heptachlor	SW-8080A	GC/ECD	0.2	ug/L	500	ug/Kg
Heptachlor Epoxide	SW-8080A	GC/ECD	0.2	ug/L	500	ug/Kg
alpha-BHC	SW-8080A	GC/ECD	0.2	ug/L	500	ug/Kg
beta-BHC	SW-8080A	. GC/ECD	0.2	ug/L	500	ug/Kg
gamma-BHC (Lindane)	SW-8080A	GC/ECD	0.2	ug/L	500	ug/Kg

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Parameter	Method Method Pescripti	Method Description	Reporting Limit		
r ar ame ter		-	Aque	ous	Non-Aqueous
delta-BHC	SW-8080A	GC/ECD	0.2	ug/L	500 ug/Kg
Methoxychlor	A0808-W2	GC/ECD	0.2	ug/L	500 ug/Kg
Toxaphene	SW-8080A	GC/ECD	0.5	ug/L	500 ug/Kg
PCB-1016	SW-8080A	GC/ECD	0.2	ug/L	500 ug/Kg
PCB-1221	SW-8080A	GC/ECD	0.2	ug/L	500 ug/Kg
PCB-1232	SW-8080A	GC/ECD	0.2	ug/L	500 ug/Kg
PCB-1242	SW-8080A	GC/ECD	0.2	ug/L	500 ug/Kg
PCB-1248	SW-8080A	GC/ECD	0.2	ug/L	500 ug/Kg
PCB-1254	SW-8080A	GC/ECD	0.2	ug/L	500 ug/Kg
PCB-1260	SW-8080A	GC/ECD	0.2	ug/L	500 ug/Kg
- Glycols					
Ethylene Glycol	SW-8015 Modified	GC	1.0	mg/L	NA
Propylene Glycol	SW-8015 Modified	GC	1.0	mg/L	NA
Diethlylene Glycol	SW-8015 Modified	GC	1.0	mg/L	NA
- Alcohols					
Methanol	SW-8015 Modified	GC	5.0	mg/L	NA
Acetonitrile	SW-8015 Modified	GC	2.0	mg/L	NA
1,4-Dioxane	SW-8015 Modified	GC	5.0	mg/L	NA
Isobutanol	SW-8015 Modified	GC	3.0	mg/L	NA
n-Butanol	SW-8015 Modified	GC	5.0	mg/L	NA
- Volatiles					
Benzene	SW-8021B/SW-8020A	GC	1.0	ug/L	5.0 ug/Kg
Ethylbenzene	SW-8021B/SW-8020A	GC	1.0	ug/L	5.0 ug/Kg
Toluene	SW-8021B/SW-8020A	GC	1.0	ug/L	5.0 ug/Kg
m&p-Xylene	SW-8021B/SW-8020A	GC	1.0	ug/L	5.0 ug/Kg

Parameter	Method Reference	Method Description	Reporting Limit		
rai alletei	Reference		Aque	ous	Non-Aqueous
o-Xylene	SW-8021B/SW-8020A	GC	1.0	ug/L	5.0 ug/Kg
Methyl-tert-butyl-ether	SW-8021B/SW-8020A	GC	1.0	ug/L	5.0 ug/Kg
C - Polynuclear Aromatic Hydro	ocarbons		· •		
Napthalene	SW-8310	HPLC	2.0	ug/L	200 ug/Kg
Acenaphthylene	SW-8310	HPLC	1.0	ug/L	100 ug/Kg
Acenaphthene	sw-8310	HPLC	1.0	ug/L	100 ug/Kg
Fluorene	sw-8310	HPLC	1.0	ug/L	100 ug/Kg
Phenanthrene	sw-8310	HPLC	1.0	ug/L	100 ug/Kg
Anthracene	sw-8310	HPLC	2.0	ug/L	100 ug/Kg
Fluoranthene	sw-8310	HPLC	0.2	ug/L	20 ug/Kg
Pyrene	SW-8310	HPLC	0.2	ug/L	20 ug/Kg
Benzo(a)anthracene	SW-8310	HPLC	0.2	ug/L	20 ug/Kg
Chrysene	SW-8310	HPLC	0.2	ug/L	20 ug/Kg
Benzo(b)fluoranthene	sw-8310	HPLC	0.2	ug/L	20 ug/Kg
Benzo(k)fluoranthene	SW-8310	HPLC	0.2	ug/L	20 ug/Kg
Benzo(a)pyrene	sw-8310	HPLC	0.2	ug/L	20 ug/Kg
Dibenz(ah)anthracene	sw-8310	HPLC	0.2	ug/L	20 ug/Kg
Benzo(ghi)perylene	sw-8310	HPLC	0.2	ug/L	20 ug/Kg
Indeno(1,2,3-cd)pyrene	s⊮-8310	HPLC	0.2	ug/L	20 ug/Kg
tal Petroleum Hydrocarbons	EPA 418.1	IR	2.0	mg/L	10 mg/Kg
tal Petroleum Hydrocarbons Diesel Range Organics)	SW-8015B	GC	0.1	mg/L	4.0 mg/Ks
tal Petroleum Hydrocarbons Gasoline Range Organics)	sw-8015B/sw-8015A Modified	GC	0.1	mg/L	0.5 mg/Ks

35. Sair Table 9.3. Analytical Methods and Reporting Limits - NPDES

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Parameter	Method Reference	Method Description	Reportin	g Limit mit
rai ametei	Reference	·	Aqueous	Non-Aqueous
Alkalinity	EPA-310.1/SM-2320B	Titration	10. mg/L	NA
Biochemical Oxygen Demand (BOD)	EPA-405.1/SM-5210B	DO Probe	4. mg/L	NA
Carbonaceous BOD (CBOD)	SM-5210B	DO Probe	4. mg/L	NA
Chemical Oxygen Demand (COD)	EPA-410.4/Hach 8000	Spectrophotometric	10. mg/l	NA
Chloride	SM-4500Cl-C	Mercuric Nitrate	5. mg/L	NA
Total Residual Chlorine	SM-4500Cl-G	DPD Colorimetric	0.1	NA
Coliform, Fecal	SM-9222 D	Membrane Filter	•	
Coliform, Total	MMO-MUG	Colilert/Colisure	Presence	/Absence
Coliform, E. Coli	MMO-MUG	Colilert/Colisure	Presence	/Absence
Color	SM-2120 B	Platinum Cobalt Units	1. C.U.	NA
Conductivity	EPA -120.1/SM-2510 B	umhos 25 degrees C	1. umhos	NA
Cyanide, Amenable	EPA-335.1/SM-4500CN-E,G	Mod. Spectrophotometric	0.005 mg/L	0.125 mg/Kg
Cyanide, Free	SM-4500CN-I	Mod. Spectrophotometric	0.005 mg/L	0.125 mg/Kg
Cyanide, Total	EPA-335.2/SM-4500CN-E	Mod. Spectrophotometric	0.005 mg/L	0.125 mg/Kg
Density	SM-2710 F			
Fluoride, Dístilled	EPA-340.1,.2/SM-4500F,B,C	Ion-Selective Electrode	0.2 mg/L	NA
Hardness, Total (CaCO3)	EPA-130.2/SM-2340C	Titration, EDTA	5.0 mg/L	NA
Hexavalent Chromium	SM-3500-Cr D	Colorimetric	0.010 mg/L	5.0 mg/Kg
Nitrogen, Ammonia Free (Direct)	EPA-350.1/SM-4500NH3	Automated Phenate	0.05 mg/L	NA
Distilled	EPA-350.1/SM-4500NH3	Automated Phenate	0.3 mg/L	30. mg/Kg
Nitrogen, Kjeldahl	EPA-350.1/SM-4500NH3	Automated Phenate	0.5 mg/L	150 mg/Kg
Nitrogen, Nitrate	EPA-353.2/SM-4500-NO3 F	Automated Cd Reduction	0.02 mg/L	0.20 mg/Kg
Nitrogen, Nitrite	EPA-353.2/SM-4500-NO3 F	Automated Cd Reduction	0.02 mg/L	0.20 mg/Kg
Dil & Grease	EPA-413.1/SM-55208,D	Gravimetric	5.0 mg/L	NA
Odor	SM 2150B		NA	NA

Parameter	Method Reference	Method Description	R	eporting Lin	g Limit mit	
			Aqueo	us	Non-	Aqueous
Oxygen, Dissolved	EPA-360.1/SM-4500-0 G.	Membrane Electrode	1.	mg/L	NA	
рН	EPA-150.1/SM-4500H-B	Potentiometric	NA		NA	
Phenols	EPA-420.1	Colorimetric	0.010	mg/L	0.25	mg/Kg
Phosphorus, Ortho	EPA-365.2/SM-4500P-E	Spectrophotometric	0.10	mg/L	NA	
Phosphorus, Total	EPA-365.2/SM-4500P-E	Spectrophotometric	0.10	mg/L	20.	mg/Kg
Sulfate	EPA-375.4	Turbidimetric	5.	mg/L	NA	
Sulfide, Total	EPA-376.1/SM-4500-S2 E	Titration	1.	mg/L	NA	
Sulfite	EPA-377.1	Titration	1.	mg/L	NA	
Surfactants (MBAS)	EPA-425.1/SM-5540-C	Colorimetric	0.030	mg/L	NA	
Total Dissolved Solids	EPA-160.1/SM-2540C	Gravimetric, 180°C	50.	mg/L	NA	
Total Suspended Solids	EPA-160.2/SM-2540D	Gravimetric, 103-105°C	3.	mg/L	NA	
Total Solids	EPA-160.3/SM-2540B	Gravimetric, 103-105°C	50.	mg/L	NA	
Total Volatile Solids	EPA-160.4	Gravimetric, 550°C	0.01	%	NA	
Total Organic Carbon (TOC)	SM-5310 B	Oxidation	1.0	mg/L	NA	
Total Petroleum HydroCarbons	EPA-418.1	Solvent extraction, IR	2.0	mg/L	10.	mg/Kg
Turbidity	EPA-180.1	Nephelometric	1.0	NTU	NA	
detals						
Aluminum (Al)	EPA-200.7 EPA-200.8	ICP ICP-MS	0.10 0.050		5.0 50.0	mg/Kg mg/Kg
Antimony (Sb)	EPA-200.7	ICP	0.10	mg/L	5.0	mg/Kg
	EPA-200.8 EPA-204.2	ICP-MS GFAA	0.001 0.020		1.0 1.0	mg/Kg mg/Kg
Arsenic (As)	EPA-200.7	ICP	0.10		5.0	mg/Kg
	EPA-200.8	ICP-MS	0.005	mg/L	5.0	mg/Kg
	EPA-206.2	GFAA	0.005	mg/L	0.25	mg/Kg
Barium (Ba)	EPA-200.7	ICP	0.020	mg/L	1.0	mg/Kg
	EPA-200.8	ICP-MS	0.005	mg/L	5.0	mg/Kg

4.1

Table 9.3.	Continued				
- 080	<del></del>				
	Method	Method	Reportir	ng Limit	
Parameter	Reference	Description		imit	
		<b>1</b>	Aqueous	Non-	Aqueous
Beryllium (Be)	EPA-200.7	ICP	0.005 mg/L	0.25	mg/Kg
	EPA-200.8	ICP-MS	0.001 mg/L	1.0	mg/Kg
**	EPA-210.2	GFAA	0.001 mg/L	0.05	mg/Kg
Boron (B)	EPA-200.7	1CP	0.050 mg/L	2.5	mg/Kg
Cadmium (Cd)	EPA-200.7	ICP	-0.030 mg/L	1.5	mg/Kg
	EPA-200.8	ICP-MS	0.001 mg/L	1.0	mg/Kg
	EPA-213.2	GFAA	0.001 mg/L	0.05	mg/Kg
Calcium (Ca)	EPA-200.7	ICP	1.0 mg/L	50.0	mg/Kg
Chromium (Cr)	EPA-200.7	ICP	0.040 mg/L	2.0	mg/Kg
	EPA-200.8	ICP-MS	0.002  mg/L	2.0	mg/Kg
	EPA-218.2	GFAA	0.002 mg/L	0.1	mg/Kg
Cobalt (Co)	EPA-200.7	ICP	0.020 mg/L	1.0	mg/Kg
	EPA-200.8	ICP-MS	0.005 mg/L	5.0	mg/Kg
	EPA-219.2	GFAA	0.005 mg/L	0.25	mg/Kg
Copper (Cu)	EPA-200.7	ICP	0.020 mg/L	1.0	mg/Kg
	EPA-200.8	ICP-MS	0.005 mg/L	5.0	mg/Kg
Iron (Fe)	EPA-200.7	ICP	0.10 mg/L	5.0	mg/Kg
Lead (Pb)	EPA-200.7	ICP	0.080 mg/L	4.0	mg/Kg
	EPA-200.8	ICP-MS	0.001 mg/L	1.0	mg/Kg
	EPA-239.2	GFAA	0.005 mg/L	0.25	mg/Kg
Magnesium (Mg)	EPA-200.7	ICP	1.0 mg/L	50.0	mg/Kg
Manganese (Mn)	EPA-200.7	ICP	0.010 mg/L	0.50	mg/Kg
	EPA-200.8	ICP-MS	0.010 mg/L	10.0	mg/Kg
Mercury (Hg)	EPA 245.1/245.5	Automated Cold Vapor	0.0002 mg/L	0.01	mg/Kg
Molybdenum (Mo)	EPA-200.7	ICP	0.020 mg/L	1.0	mg/Kg
	EPA-200.8	ICP-MS	0.001 mg/L	1.0	mg/Kg
Nickel (Ni)	EPA-200.7	ICP	0.010 mg/L	0.50	mg/Kg
	EPA-200.8	ICP-MS	0.005 mg/L	5.0	mg/Kg
Potassium (K)	EPA-200.7	ICP	1.0 mg/L	50.0	mg/Kg
Selenium (Se)	EPA-200.7	ICP	0.10 mg/L	5.0	mg/Kg
	EPA-200.8	ICP-MS	0.005 mg/L	5.0	mg/Kg
	EPA-270.2	´ GFAA	0.005 mg/L	0.25	mg/Kg

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Parameter	Method Reference	Method Description	Repor	ting Limit Limit		
		<i>h</i>	Aqueous		Non-Aqueous	
Silver (Ag)	EPA-200.7	ICP	0.040 mg/	L 2.0	mg/Kg	
, <del>a</del>	EPA-200.8	ICP-MS	0.0005 mg/	L 0.5	mg/Kg	
	EPA-272.2	GFAA	0.001 mg/	L 0.05	mg/Kg	
Sodium (Na)	EPA-200.7	ICP	1.0 mg/	L 50.0	mg/Kg	
Strontium (Sr)	EPA-200.7	ICP	0.10 mg/	L 5.0	mg/Kg	
Thallium (Tl)	EPA-200.7	ICP	0.50 mg/	L 25	mg/Kg	
	EPA-200.8	ICP-MS	0.001 mg/	L 1.0	mg/Kg	
	EPA-279.2	GFAA	0.010 mg/	L 0.5	mg/Kg	
Tin (Sn)	EPA-200.7	ICP	2.0 mg/	L 100	mg/Kg	
Titanium (Ti)	EPA-200.7	ICP	0.020 mg/	L 1.0	mg/Kg	
Vanadium (V)	EPA-200.7	ICP	0.050 mg/	L 2.5	mg/Kg	
	EPA-200.8	ICP-MS	0.005 mg/	L 5.0	mg/Kg	
Zinc (Zn)	EPA-200.7	1CP	0.050 mg/	L 2.5	mg/Kg	
	EPA-200.8	ICP-MS	0.050 mg/		mg/Kg	
rganics - Volatiles						
Acetone	EPA-624	GC/MS	20 ug/	L 5.0	mg/Kg	
Acrolein (Screen)	EPA-624	GC/MS	50 ug/	L 12.5	mg/Kg	
Acrylonitrile (Screen)	EPA-624	GC/MS	50 ug/	L 5.0	mg/kg	
Benzene	EPA-624	GC/MS	1.0 ug/	L 0.25	mg/Kg	
Bromodichloromethane	EPA-624	GC/MS	1.0 ug/	L 0.25	mg/Kg	
Bromoform	EPA-624	GC/MS	1.0 ug/	L 0.25	mg/Kg	
Bromomethane	EPA-624	GC/MS	1.0 ug/	L 0.25	mg/Kg	
2-Butanone (MEK)	EPA-624	GC/MS	10.0 ug/	L 2.5	mg/Kg	
Carbon Disulfide	EPA-624	GC/MS	1.0 ug/	L 0.25	mg/Kg	
Carbon Tetrachloride	EPA-624	GC/MS	1.0 ug/	L 0.25	mg/Kg	
Chlorobenzene	EPA-624	GC/MS	1.0 ug/	L 0.25	mg/Kg	
Chloroethane	EPA-624	GC/MS	5.0 ug/	L 0.25	mg/Kg	
2-Chloroethylvinyl ether	EPA-624	GC/MS	1.0 ug/	L 0.25	mg/Kg	

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Parameter	Method Reference	Method Description	Reporting Limit Limit			
			Aqueo	us	Non-/	Aqueous
Chloroform	EPA-624	GC/MS	1.0	ug/L	0.25	mg/Kg
Chloromethane	EPA-624	GC/MS	10.0	ug/L	2.5	mg/Kg
Dibromochloromethane	EPA-624	GC/MS	1.0	ug/L	0.25	mg/Kg
1,2-Dichlorobenzene	EPA-624	GC/MS	1.0	ug/L	0.25	mg/Kg
1,3-Dichlorobenzene	EPA-624	GC/MS	1.0	ug/L	0.25	mg/Kg
1,4-Dichlorobenzene	EPA-624	GC/MS	1.0	ug/L	0.25	mg/Kg
1,1-Dichloroethane	EPA-624	GC/MS	1.0	ug/L	0.25	mg/Kg
1,2-Dichl <i>o</i> roethane	EPA-624	GC/MS	1.0	ug/L	0.25	mg/Kg
1,1-Dichloroethene	EPA-624	GC/MS	1.0	ug/L	0.25	mg/Kg
cis-1,2,Dichloroethene	EPA-624	GC/MS	1.0	ug/L	0.25	mg/Kg
trans-1,2-Dichloroethene	EPA-624	GC/MS	1.0	ug/L	0.25	mg/Kg
1,2-Dichloropropane	EPA-624	GC/MS	1.0	ug/L	0.25	mg/Kg
cis-1,3-Dichloropropene	EPA-624	GC/MS	1.0	ug/L	0.25	mg/Kg
trans-1,3-Dichloropropene	EPA-624	GC/MS	1.0	ug/L	0.25	mg/Kg
Ethyl benzene	EPA-624	GC/MS	1.0	ug/L	0.25	mg/Kg
Methylene Chloride	EPA-624	GC/MS	10.0	ug/L	0.25	mg/Kg
4-Methyl-2-pentanone (MIBK)	EPA-624	GC/MS	10.0	ug/L	0.25	mg/Kg
Styrene	EPA-624	GC/MS	1.0	ug/L	0.25	mg/Kg
1,1,2,2-Tetrachloroethane	EPA-624	GC/MS	1.0	ug/L	0.25	mg/Kg
1,2,4-Trimethylbenzene	EPA-624	GC/MS	1.0	ug/L	0.25	mg/Kg
1,3,5-Trimethylbenzene	EPA-624	GC/MS	1.0	ug/L	0.25	mg/Kg
Tetrachloroethene	EPA-624	GC/MS	1.0	ug/L	0.25	mg/Kg
Toluene	EPA-624	GC/MS	1.0	ug/L	0.25	mg/Kg
1,1,1-Trichloroethane	EPA-624	· GC/MS	1.0	ug/L	0.25	mg/Kg
1,1,2-Trichloroethane	EPA-624	GC/MS	1.0	ug/L	0.25	mg/Kg

Parameter	Method Reference	Method Description	Reporting Limit Limit			
		<i>p</i>	Aqueo			Aqueous
Trichloroethene	EPA-624	GC/MS	1.0	ug/L	0.25	mg/Kg
Vinyl Chloride	EPA-624	GC/MS	2.0	ug/L	0.25	mg/Kg
Xylenes, total	EPA-624	GC/MS	1.0	ug/L	0.25	mg/Kg
rganics - Semi-Volatiles			-			
Acenaphthene	EPA-625	GC/MS	5.	ug/L	165	ug/Kg
Acenaphthylene	EPA-625	GC/MS	5.	ug/L	165	ug/Kg
Anthracene	EPA-625	GC/MS	5.	ug/L	165	ug/Kg
Benzidine	EPA-625	GC/MS	50.	ug/L	1,650	ug/Kg
Benzo(a)anthracene	EPA-625	GC/MS	5.	ug/L	165	ug/Kg
Benzo(b)fluoranthene	EPA-625	GC/MS	10.	ug/L	330	ug/Kg
Benzo(k)fluoranthene	EPA-625	GC/MS	10.	ug/L	330	ug/Kg
Benzo(a)pyrene	EPA-625	GC/MS	5.	ug/L	165	ug/Kg
Benzo(g,h,i)perylene	EPA-625	GC/MS	5.	ug/L	165	ug/Kg
Benzyl butyl phthalate	EPA-625	GC/MS	5.	ug/L	165	ug/Kg
Bis(2-chloroethyl)ether	EPA-625	GC/MS	5.	ug/L	165	ug/Kg
Bis(2-chloroethoxy)methane	EPA-625	GC/MS	5.	ug/L	165	ug/Kg
Bis(2-ethylhexyl)phthalate	EPA-625	GC/MS	5.	ug/L	165	ug/Kg
Bis(2-chloroisopropyl)ether	EPA-625	GC/MS	5.	ug/L	165	ug/Kg
4-Bromophenyl phenyl ether	EPA-625	GC/MS	5.	ug/L	165	ug/Kg
2-Chloronaphthalene	EPA-625	GC/MS	5.	ug/L	165	ug/Kg
4-Chlorophenyl phenyl ether	EPA-625	GC/MS	5.	ug/L	165	ug/Kg
Chrysene	EPA-625	GC/MS	5.	ug/L	165	ug/Kg
Dibenzo(a,h)anthracene	EPA-625	GC/MS	5.	ug/L	165	ug/Kg
Di-n-butylphthalate	EPA-625	· GC/MS	5.	ug/L	165	ug/Kg
1,2-Dichlorobenzene	EPA-625	GC/MS	5.	ug/L	165	ug/Kg

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Parameter  1,3-Dichlorobenzene	Method Reference	Method Description	Reporting Limit Limit			
			Aque	ous	Non-	Aqueous
	EPA-625	GC/MS	5.	ug/L	165	ug/Kg
1,4-Dichlorobenzene	EPA-625	GC/MS	5.	ug/L	165	ug/Kg
3,3-Dichlorobenzidine	EPA-625	GC/MS	50.	ug/L	1,650	ug/Kg
1,2-Diphenylhydrazine	EPA-625	GC/MS	5.	ug/L	165	ug/Kg
Diethyl phthalate	EPA-625	GC/MS	5.	ug/L	165	ug/Kg
Dimethyl phthalate	EPA-625	GC/MS	5.	ug/L	165	ug/Kg
2,4-Dinitrotoluene	EPA-625	GC/MS	5.	ug/L	165	ug/Kg
2,6-Dinitrotoluene	EPA-625	GC/MS	5.	ug/L	165	ug/Kg
Di-n-octylphthalate	EPA-625	GC/MS	5.	ug/L	165	ug/Kg
Fluoranthene	EPA-625	GC/MS	5.	ug/L	165	ug/Kg
Fluorene	EPA-625	GC/MS	5.	ug/L	165	ug/Kg
Hexachlorobenzene	EPA-625	GC/MS	5.	ug/L	165	ug/Kg
<b>Hexachlorobutadiene</b>	EPA-625	GC/MS	5.	ug/L	165	ug/Kg
Hexachlorocyclopentadiene	EPA-625	GC/MS	20.	ug/L	660	ug/Kg
Hexachloroethane	EPA-625	GC/MS	5.	ug/L	165	ug/Kg
Indeno(1,2,3-cd)pyrene	EPA-625	GC/MS	5.	ug/L	165	ug/Kg
Isophorone	EPA-625	GC/MS	5.	ug/L	165	ug/Kg
Naphthalene	EPA-625	GC/MS	5.	ug/L	165	ug/Kg
Nitrobenzene	EPA-625	GC/MS	5.	ug/L	165	ug/Kg
N-Nitrosodimethylamine	EPA-625	GC/MS	5.	ug/L	165	ug/Kg
N-Nitrosodiphenylamine	EPA-625	GC/MS	5.	ug/L	165	ug/Kg
N-Nitroso-di-n-propylamine	EPA-625	GC/MS	5.	ug/L	165	ug/Kg
Phenanthrene	EPA-625	GC/MS	5.	ug/L	165	ug/Kg
Pyrene	EPA-625	· GC/MS	5.	ug/L	165	ug/Kg
1,2,4-Trichlorobenzene	EPA-625	GC/MS	5.	ug/L	165	ug/Kg

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Parameter	Method Reference	Method Description	Reporting Limit Limit			
			Aqueo	ous	Non-	Aqueous
4-Chloro-3-methylphenol	EPA-625	GC/MS	10.	ug/L	330	ug/Kg
2-Chlorophenol	EPA-625	GC/MS	10.	ug/L	330	ug/Kg
2,4-Dichlorophenol	EPA-625	GC/MS	10.	ug/L	330	ug/Kg
2,4-Dimethylphenol	EPA-625	GC/MS	10.	ug/L	330	ug/Kg
2,4-Dinitrophenol	EPA-625	GC/MS	10.	ug/L	330	ug/Kg
2-Methyl-4,6-dinitrophenol	EPA-625	GC/MS	10.	ug/L	330	ug/Kg
2-Nitrophenol	EPA-625	GC/MS	10.	ug/L	330	ug/Kg
4-Nitrophenol	EPA-625	GC/MS	10.	ug/L	330	ug/Kg
Pentachlorophenol	EPA-625	GC/MS	10.	ug/L	330	ug/Kg
Phenol	EPA-625	GC/MS	10.	ug/L	330	ug/Kg
2,4,6-Trichlorophenol	EPA-625	GC/MS	10.	ug/L	330	ug/Kg
2-Methylphenol	EPA-625	GC/MS	10.	ug/L	330	ug/Kg
4-Methylphenol	EPA-625	GC/MS	10.	ug/L	330	ug/Kg
organics - Pesticides/PCBs						
Aldrin	EPA 608	GC/ECD	0.2	ug/L	500	ug/Kg
Chlordane	EPA 608	GC/ECD	0.2	ug/L	500	ug/Kg
Dieldrin	EPA 608	GC/ECD	0.2	ug/L	500	ug/Kg
4,4'-DDD	EPA 608	GC/ECD	0.2	ug/L	500	ug/Kg
4,4'-DDE	EPA 608	GC/ECD	0.2	ug/L	500	ug/Kg
4,4'-DDT	EPA 608	GC/ECD	0.2	ug/L	500	ug/Kg
Endosulfan I	EPA 608	GC/ECD	0.2	ug/L	500	ug/Kg
Endosulfan II	EPA 608	GC/ECD	0.2	ug/L	500	ug/Kg
Endosulfan Sulfate	EPA 608	GC/ECD	0.2	ug/L	500	ug/Kg
Endrin	EPA 608	· GC/ECD	0.2	ug/L	500	ug/Kg
Endrin Aldehyde	EPA 608	GC/ECD	0.2	ug/L	500	ug/Kg

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Parameter	Method Reference	Method Description	Reporting Lim Aqueous	
Endrin Ketone	EPA 608	GC/ECD	0.2 ug/L	500 ug/Kg
Heptachlor	EPA 608	GC/ECD	0.2 ug/L	500 ug/Kg
Heptachlor Epoxide	EPA 608	GC/ECD	0.2 ug/L	500 ug/Kg
alpha-BHC	EPA 608	GC/ECD	0.2 ug/L	500 ug/Kg
beta-BHC	EPA 608	GC/ECD	0.2 ug/L	500 ug/Kg
gamma-BHC (Lindane)	EPA 608	GC/ECD	0.2 ug/L	500 ug/Kg
delta-BHC	EPA 608	GC/ECD	0.2 ug/L	500 ug/Kg
Methoxychlor	EPA 608	GC/ECD	0.2 ug/L	500 ug/Kg
Toxaphene	EPA 608	GC/ECD	0.5 ug/L	500 ug/Kg
PCB-1016	EPA 608	GC/ECD	0.2 ug/L	500 ug/Kg
PCB-1221	EPA 608	GC/ECD	0.2 ug/L	500 ug/Kg
PCB-1232	EPA 608	GC/ECD	0.2 ug/L	500 ug/Kg
PCB-1242	EPA 608	GC/ECD	0.2 ug/L	500 ug/Kg
PCB-1248	EPA 608	GC/ECD	0.2 ug/L	500 ug/Kg
PCB-1254	EPA 608	GC/ECD	0.2 ug/L	500 ug/Kg
PCB-1260	EPA 608	GC/ECD	0.2 ug/L	500 ug/Kg
rganics - GC Volatiles				
1,2-Dibromo-3-chloropropane	EPA-504.1	GC/ECD	0.02 ug/L	NA
Ethylene Dibromide	EPA-504.1	GC/ECD	0.02 ug/Ł	NA

Table 9.4. Analytical Methods and Reporting Limits - Ohio VAP

. (2)						
Parameter	Method Reference	Method Description	Reporting Limit			
		•.	Aqueous	Non-Aqueous		
Wet Chemistry						
Cyanide, Total	EPA 335.2 CLP M	Spectrophotometric	0.005 mg/L	0.125 mg/Kg		
Hexavalent Chromium	sw-7196A	Colorimetric	0.010 mg/L	5.0 mg/Kg		
Phosphorus, Ortho	EPA-365.2	Spectrophotometric	0.10 mg/L	20. mg/Kg		
Phosphorus, Total	EPA-365.2	Spectrophotometric	0.10 mg/L	20. mg/Kg		
Metals						
Aluminum (Al)	SW-6010A	ICP	0.10 mg/L	5.0 mg/Kg		
	sw-6020	ICP-MS	0.050 mg/L	50.0 mg/Kg		
Antimony (Sb)	SW-6010A	ICP	0.10 mg/L	5.0 mg/Kg		
The timothy (OD)	SW-6020	ICP-MS	0.001 mg/L	1.0 mg/Kg		
	sw-7041	GFAA	0.020 mg/L	1.0 mg/Kg		
Arsenic (As)	SW-6010A	ICP	0.10 mg/L	5.0 mg/Kg		
	sw-6020	ICP-MS	0.005 mg/L	5.0 mg/Kg		
	SW-7060A	GFAA	0.005 mg/L	0.25 mg/Kg		
Barium (Ba)	SW-6010A	ICP	0.020 mg/L	1.0 mg/Kg		
	sW-6020	ICP-MS	0.005 mg/L	5.0 mg/Kg		
Beryllium (Be)	SW-6010A	ICP	0.005 mg/L	0.25 mg/Kg		
	sw-6020	ICP-MS	0.001 mg/L	1.0 mg/Kg		
	sw-7091	GFAA .	0.001 mg/L	0.05 mg/Kg		
Cadmium (Cd)	SW-6010A	ICP	0.030 mg/L	1.5 mg/Kg		
	sw-6020	ICP-MS	0.001  mg/L	1.0 mg/Kg		
	SW-7131A	GFAA	0.001 mg/L	0.05 mg/Kg		
Calcium (Ca)	SW-6010A	1CP	1.0 mg/L	50.0 mg/Kg		
Chromium (Cr)	SW-6010A	ICP	0.040 mg/L	2.0 mg/Kg		
	sw-6020	ICP-MS	0.002 mg/L	2.0 mg/Kg		
	sw-7191	GFAA	0.002 mg/L	0.1 mg/Kg		
Cobalt (Co)	SW-6010A	ICP	0.020 mg/L	1.0 mg/Kg		
	SW-6020	ICP-MS	0.005 mg/L	5.0 mg/Kg		
Copper (Cu)	SW-6010A	ICP	0.020 mg/L	1.0 mg/Kg		
	sw-6020	ICP-MS	0.005 mg/L	5.0 mg/Kg		
Iron (Fe)	SW-6010A	1CP	0.10 mg/L	5.0 mg/Kg		

	Method	Method	1	Reporti	ng Limit	
Parameter	Reference	Description	Aque	ous	Non	-Aqueous
Lead (Pb)	SW-6010A	ICP	0.080	mg/L	4.0	mg/Kg
	sw-6020	ICP-MS	0.001	mg/L	1.0	mg/Kg
*	sw-7421	GFAA	0.005	mg/L	0.25	mg/Kg
Manganese (Mn)	SW-6010A	ICP	0.010	mg/L	0.50	mg/Kg
	sw-6020	ICP-MS	0.010	mg/L	10.0	mg/Kg
Mercury (Hg)	sw-7470a/sw-7471a	Automated Cold Vapor	0.0002	mg/L	0.01	mg/Kg
Nickel (Ni)	SW-6010A	ICP	0.010	mg/L	0.5	mg/Kg
	SW-6020	ICP-MS	0.005	mg/L	5.0	mg/Kg
Potassium (K)	SW-6010A	ICP	1.0	mg/L	50.0	mg/Kg
Selenium (Se)	SW-6010A	ICP	0.10	mg/L	5.0	mg/Kg
	sw-7740	GFAA	0.005	mg/L	0.25	mg/Kg
Silver (Ag)	SW-6010A	ICP	0.040	mg/L	2.0	mg/Kg
	sw-6020	ICP-MS	0.0005		0.5	mg/Kg
	sw-7761	GFAA	0.001	mg/L	0.05	mg/Kg
Sodium (Na)	SW-6010A	ICP	1.0	mg/L	50.0	mg/Kg
Thallium (Tl)	SW-6010A	1CP	0.50			mg/Kg
	sw-6020	I CP-MS	0.001			mg/Kg
	sw-7841	GFAA	0.010	mg/L	0.5	mg/Kg
Vanadium (V)	SW-6010A	ICP	0.050	mg/L	2.5	mg/Kg
	EPA-200.8	ICP-MS	0.005	mg/L		mg/Kg
Zinc (Zn)	SW-6010A	ICP	0.050	mg/L	2.5	mg/Kg
	SW-6020	1CP-MS	0.050	mg/L	50	mg/Kg
Organics - Volatile Compounds						
Acetone	SW-8260A	GC/MS	20	ug/L	100	ug/Kg
Acrolein	SW-8260A	GC/MS	50	ug/L	50	ug/Kg
Acrylonitrile	SW-8260A	GC/MS	50	ug/L	50	ug/Kg
Allyl chloride	SW-8260A	GC/MS	5.0	ug/L	5.0	ug/Kg
Benzene	SW-8260A	GC/MS	1.0	ug/L	5.0	ug/Kg
	EPA 524.2	GC/MS	0.5	ug/L	N/	A
Bromobenzene	SW-8260A	GC/MS	1.0	ug/L	5.0	ug/Kg
	EPA 524.2	GC/MS	0.5	ug/L	N/	4
Bromochloromethane	SW-8260A	GC/MS	1.0	ug/L		ug/Kg
	EPA 524.2	GC/MS	0.5	ug/L	N/	A

2.

Table 9.4. Continued

Parameter  Bromodichloromethane	Method	Method		Reporti	ng Limit
	Reference	Description	upA	eous	Non-Aqueou
	SW-8260A	GC/MS	1.0	ug/L	5.0 ug/Kg
*	EPA 524.2	GC/MS	0.5	ug/L	NA
Bromoform	SW-8260A	GC/MS	1.0	ug/L	5.0 ug/Kg
	EPA 524.2	GC/MS	0.5	ug/L	NA
Bromomethane	SW-8260A	GC/MS	5.0	ug/L	5.0 ug/Kg
	EPA 524.2	GC/MS	0.5	ug/L	NA
n-Butylbenzene	SW-8260A	GC/MS	1.0	ug/L	5.0 ug/Kg
	EPA 524.2	GC/MS	0.5	ug/L	NA
tert-Butylbenzene	SW-8260A	GC/MS	· 1.0	ug/L	5.0 ug/Kg
·	EPA 524.2	GC/MS	0.5	ug/L	NA
sec-Butylbenzene	SW-8260A	GC/MS	1.0	ug/L	5.0 ug/Kg
,	EPA 524.2	GC/MS	0.5	ug/L	NA
2-Butanone (MEK)	SW-8260A	GC/MS	20	ug/L	100 ug/Kg
Carbon Disulfide	SW-8260A	GC/MS	1.0	ug/L	5.0 ug/Kg
Carbon Tetrachloride	SW-8260A	GC/MS	1.0	ug/L	5.0 ug/Kg
	EPA 524.2	GC/MS	0.5	ug/L	NA
Chlorobenzene	SW-8260A	GC/MS	1.0	ug/L	5.0 ug/Kg
	EPA 524.2	GC/MS	0.5	ug/L	NA
Chloroethane	SW-8260A	GC/MS	5.0	ug/L	5.0 ug/Kg
	EPA 524.2	GC/MS	0.5	ug/L	NA
2-Chloroethylvinyl ether	SW-8260A	GC/MS	5.0	ug/L	5.0 ug/Kg
Chloroform	SW-8260A	GC/MS	1.0	ug/L	5.0 ug/Kg
	EPA 524.2	GC/MS	0.5	ug/L	NA
Chloromethane	SW-8260A	GC/MS	5.0	ug/L	10.0 ug/Kg
	EPA 524.2	GC/MS	0.5	ug/L	NA
Chloroprene	SW-8260A	GC/MS	5.0	ug/L	5.0 ug/Kg
o-Chlorotoluene	SW-8260A	GC/MS	1.0	ug/L	5.0 ug/Kg
	EPA 524.2	GC/MS	0.5	ug/L	NA
p-Chlorotoluene	SW-8260A	GC/MS	1.0	ug/L	5.0 ug/Kg
,	EPA 524.2	GC/MS	0.5	ug/L	NA NA
Dibromochloromethane	SW-8260A	GC/MS	1.0	ug/L	5.0 ug/Kg
,	EPA 524.2	GC/MS	0.5	ug/L	NA NA

Parameter	Method Reference	Method Description		Reportir	ng Limit
rarameter		<i>P</i>	Aqu	eous	Non-Aqueou
1,2-Dibromo-3-Chloropropane	SW-8260A	GC/MS	5.0	ug/L	5.0 ug/Kg
1,2-Dibromoethane	SW-8260A	GC/MS	5.0	ug/L	5.0 ug/Kg
Dibromomethane	SW-8260A	GC/MS	1.0	ug/L	5.0 ug/Kg
	EPA 524.2	GC/MS	۵.5	ug/L	NA
1,2-Dichlorobenzene	SW-8260A	GC/MS	1.0	ug/L	5.0 ug/Kg
,	EPA 524.2	GC/MS	0.5	ug/L	NA
1,3-Dichlorobenzene	SW-8260A	GC/MS	1.0	ug/L	5.0 ug/Kg
	EPA 524.2	GC/MS	0.5	ug/L	NA
1,4-Dichlorobenzene	SW-8260A	GC/MS	1.0	ug/L	5.0 ug/Kg
	EPA 524.2	GC/MS	0.5	ug/L	NA
trans-1,4-Dichloro-2-butene	SW-8260A	GC/MS	5.0	ug/L	5.0 ug/Kg
Dichlorodifluoromethane	SW-8260A	GC/MS	1.0	ug/L	5.0 ug/Kg
1,1-Dichloroethane	SW-8260A	GC/MS	1.0	ug/L	5.0 ug/Kg
•	EPA 524.2	GC/MS	0.5	ug/L	NA
1,2-Dichloroethane	SW-8260A	GC/MS	1.0	ug/L	5.0 ug/Kg
	EPA 524.2	GC/MS	0.5	ug/L	NA
1,1-Dichloroethene	SW-8260A	GC/MS	1.0	ug/L	5.0 ug/Kg
	EPA 524.2	GC/MS	0.5	ug/L	NA
cis-1,2,Dichloroethene	SW-8260A	GC/MS	1.0	ug/L	5.0 ug/Kg
	EPA 524.2	GC/MS	0.5	ug/L	NA
trans-1,2-Dichloroethene	SW-8260A	GC/MS	1.0	ug/L	5.0 ug/Kg
·	EPA 524.2	GC/MS	0.5	ug/L	NA
1,1-Dichloropropene	SW-8260A	GC/MS	1.0	ug/L	5.0 ug/Kg
	EPA 524.2	GC/MS	0.5	ug/L	NA
1,2-Dichloropropane	SW-8260A	GC/MS	1.0	ug/L	5.0 ug/Kg
	EPA 524.2	GC/MS	0.5	ug/L	NA
cis-1,3-Dichloropropene	SW-8260A	GC/MS	1.0	ug/L	5.0 ug/Kg
· · · · · · · · · · · · · · · · · · ·	EPA 524.2	GC/MS	0.5	ug/L	NA
trans-1,3-Dichloropropene	SW-8260A	GC/MS	1.0	ug/L	5.0 ug/Kg
· · · · · · · · · · · · · · · · · · ·	EPA 524.2	GC/MS	0.5	ug/L	NA NA
1,3-Dichloropropane	SW-8260A	GC/MS	1.0	ug/L	5.0 ug/Kg
	EPA 524.2	GC/MS	0.5	ug/L	NA NA

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Table 9.4. Continued

	Method	Method :		Reportir	ng Limit
Parameter	Reference	Description	upA	eous	Non-Aqueous
2,2-Dichloropropane	SW-8260A	GC/MS	1.0	ug/L	5.0 ug/Kg
, , , , , , , , , , , , , , , , , , ,	EPA 524.2	GC/MS	0.5	ug/L	NA NA
Ethyl benzene	SW-8260A	GC/MS	1.0	ug/L	5.0 ug/Kg
	EPA 524.2	GC/MS	0.5	ug/L	NA
Fluorotrichloromethane	SW-8260A	GC/MS	1.0	ug/L	5.0 ug/Kg
	EPA 524.2	GC/MS	0.5	ug/L	NA
Hexachlorobutadiene	SW-8260A	GC/MS	5.0	ug/L	5.0 ug/Kg
	EPA 524.2	GC/MS	0.5	ug/L	NA
2-Hexanone	SW-8260A	GC/MS	10	ug/L	50 ug/Kg
lodomethane	SW-8260A	GC/MS	5.0	ug/L	5.0 ug/Kg
Isopropylbenzene	SW-8260A	GC/MS	1.0	ug/L	5.0 ug/Kg
	EPA 524.2	GC/MS	0.5	ug/L	NA
p-Isopropyltoluene	SW-8260A	GC/MS	1.0	ug/L	5.0 ug/Kg
	EPA 524.2	GC/MS	0.5	ug/L	NA
Methylene Chloride	SW-8260A	GC/MS	10	ug/L	10 ug/Kg
	EPA 524.2	GC/MS	0.5	ug/L	NA
4-Methyl-2-pentanone (MIBK)	SW-8260A	GC/MS	10	ug/L	50 ug/Kg
Naphthalene	SW-8260A	GC/MS	5.0	ug/L	5.0 ug/Kg
	EPA 524.2	GC/MS	0.5	ug/L	NA
Propionitrile	SW-8260A	GC/MS	50	ug/L	50 ug/Kg
n-Propylbenzene	SW-8260A	GC/MS	1.0	ug/L	5.0 ug/Kg
	EPA 524.2	GC/MS	0.5	ug/L	NA
Styrene	SW-8260A	GC/MS	1.0	ug/L	5.0 ug/Kg
	EPA 524.2	GC/MS	0.5	ug/L	. NA
1,1,1,2-Tetrachloroethane	SW-8260A	GC/MS	1.0	ug/L	5.0 ug/Kg
	EPA 524.2	GC/MS	0.5	ug/L	NA
1,1,2,2-Tetrachloroethane	SW-8260A	GC/MS	1.0	ug/L	5.0 ug/Kg
	EPA 524.2	GC/MS	0.5	ug/L	NA
1,2,3-Trichloropropane	SW-8260A	GC/MS	1.0	ug/L	5.0 ug/Kg
	EPA 524.2	GC/MS	0.5	ug/L	NA NA
1,2,4-Trichlorobenzene	SW-8260A	GC/MS	5.0	ug/L	5.0 ug/Kg
	EPA 524.2	GC/MS	0.5	ug/L	NA NA

	Method	Method		Reportir	ng Limit
Parameter	Reference	Description	Aqu	eous	Non-Aqueous
1,2,3-Trichlorobenzene	SW-8260A	GC/MS	5.0	ug/L	5.0 ug/Kg
	EPA 524.2	GC/MS	0.5	ug/L	NA
1,2,4-Trimethylbenzene	SW-8260A	GC/MS	1.0	ug/L	5.0 ug/Kg
	EPA 524.2	GC/MS	0.5	ug/L	NA
1,3,5-Trimethylbenzene	SW-8260A	GC/MS	1.0	ug/L	5.0 ug/Kg
	EPA 524.2	GC/MS	0.5	ug/L	NA
Tetrachloroethene	SW-8260A	GC/MS	1.0	ug/L	5.0 ug/Kg
	EPA 524.2	GC/MS	0.5	ug/L	NA
Toluene	SW-8260A	GC/MS	1.0	ug/L	5.0 ug/Kg
	EPA 524.2	GC/MS	0.5	ug/L	NA .
1,1,1-Trichloroethane	SW-8260A	GC/MS	1.0	ug/L	5.0 ug/Kg
, .	EPA 524.2	GC/MS	0.5	ug/L	NA
1,1,2-Trichloroethane	SW-8260A	GC/MS	1.0	ug/L	5.0 ug/Kg
, .	EPA 524.2	GC/MS	0.5	ug/L	NA
Trichloroethene	SW-8260A	GC/MS	1.0	ug/L	5.0 ug/Kg
	EPA 524.2	GC/MS	0.5	ug/L	NA
Vinyl Acetate	SW-8260A	GC/MS	5.0	ug/L	5.0 ug/Kg
Vinyl Chloride	SW-8260A	GC/MS	2.0	ug/L	2.0 ug/Kg
	EPA 524.2	GC/MS	0.5	ug/L	NA
o-Xylene	SW-8260A	GC/MS	1.0	ug/L	5.0 ug/Kg
	EPA 524.2	GC/MS	0.5	ug/L	NA
m & p Xylene	SW-8260A	GC/MS	1.0	ug/L	5.0 ug/Kg
	EPA 524.2	GC/MS	0.5	ug/L	NA
Xylenes, total	SW-8260A	GC/MS	1.0	ug/L	5.0 ug/Kg
	EPA 524.2	GC/MS	0.5	ug/L	NA
n-Hexane	SW-8260A	GC/MS	10	ug/L	10 ug/Kg
MI-VOLATILE COMPOUNDS					
Acenaphthene	sw-8270B	GC/MS	10	ug/L	330 ug/Kg
Acenaphthylene	SW-8270B	GC/MS	10	ug/L	330 ug/Kg
Acetophenone	SW-8270B	, GC/MS	20	ug/L	660 ug/Kg
2-Acetylaminoflourene (2-AAF)	SW-8270B	GC/MS	20	ug/L	660 ug/Kg

Parameter	Method Reference	Method Description		Report	ing Limi	t
Parameter	Reference	)	Aqu	eous	No	n-Aqueous
4-Aminobipheỳl	SW-8270B	GC/MS	20	ug/L	660	ug/Kg
Aniline	SW-8270B	GC/MS	10	ug/L	330	ug/Kg
Anthracene	SW-8270B	GC/MS	_10	ug/L	330	ug/Kg
Aramite	SW-8270B	GC/MS	15	ug/L	495	ug/Kg
Benzidine	SW-8270B	GC/MS	50	ug/L	1,650	ug/Kg
Benzo(a)anthracene	SW-8270B	GC/MS	10	ug/L	330	ug/Kg
Benzo(b)fluoranthene	SW-8270B	GC/MS	10	ug/L	330	ug/Kg
Benzo(k)fluoranthene	SW-8270B	GC/MS	10	ug/L	330	ug/Kg
Benzo(a)pyrene	SW-8270B	GC/MS	10	ug/L	330	ug/Kg
Benzo(g,h,i)perylene	SW-82708	GC/MS	10	ug/L	330	ug/Kg
Benzył alcohol	SW-8270B	GC/MS	10	ug/L	330	ug/Kg
Benzyl butyl phthalate	SW-8270B	GC/MS	10	ug/L	330	ug/Kg
Bis(2-chloroethyl)ether	SW-8270B	GC/MS	10	ug/L	330	ug/Kg
Bis(2-chloroethoxy)methane	SW-8270B	GC/MS	10	ug/L	330	ug/Kg
Bis(2-ethylhexyl)phthalate	SW-8270B	GC/MS	10	ug/L	330	ug/Kg
Bis(2-chloroisopropyl)ether	SW-8270B	GC/MS	10	ug/L	330	ug/Kg
4-Bromophenyl phenyl ether	SW-8270B	GC/MS	10	ug/L	330	ug/Kg
4-Chloroaniline	SW-8270B	GC/MS	10	ug/L	330	ug/Kg
Chlorobenzilate	SW-8270B	GC/MS	50	ug/L	1,650	ug/Kg
2-Chloronaphthalene	SW-8270B	GC/MS	10	ug/L	330	ug/Kg
4-Chlorophenyl phenyl ether	SW-8270B	GC/MS	10	ug/L	330	ug/Kg
Chrysene	sw-8270B	GC/MS	10	ug/L	330	ug/Kg
Diallate	sw-8270B	GC/MS	30	ug/L	990	ug/Kg
Dibenzo(a,h)anthracene	SW-8270B	GC/MS	10	ug/L	330	ug/Kg
Dibenzofuran	SW-8270B	GC/MS	10	ug/L	330	ug/Kg

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Table 9.4. Continued

Davanana	Method	Method		Report	ing Limi	t
Parameter	Reference	Description	Aqu	ieous	No	n-Aqueous
Di-n-butylphthalate	SW-8270B	GC/MS	10	ug/L	330	ug/Kg
1,2-Dichlorobenzene	SW-8270B	GC/MS	10	ug/L	330	ug/Kg
1,3-Dichlorobenzene	SW-8270B	GC/MS	10	ug/L	330	ug/Kg
1,4-Dichlorobenzene	sw-8270B	GC/MS	10	ug/L	330	ug/Kg
3,3-Dichlorobenzidine	SW-8270B	GC/MS	50	ug/L	1,650	ug/Kg
Diethyl phthalate	SW-8270B	GC/MS	10	ug/L	330	ug/Kg
Dimethoate	SW-8270B	GC/MS	100	ug/L	3,300	ug/Kg
p-(Dimethylamino)-azobenzene	SW-8270B	GC/MS	50	ug/L	1,650	ug/Kg
7,12-Dimethylbenz(a)anthracene	SW-8270B	GC/MS	20	ug/L	660	ug/Kg
3,3'-Dimethylbenzidine	SW-8270B	GC/MS	20	ug/L	660	ug/Kg
a,a-Dimethyl-phenethylamino	SW-8270B	GC/MS	50	ug/L	1,650	ug/Kg
Dimethyl phthalate	S₩-8270B	GC/MS	10	ug/L	330	ug/Kg
2,4-Dinitrotoluene	SW-8270B	GC/MS	10	ug/L	330	ug/Kg
2,6-Dinitrotoluene	SW-8270B	GC/MS	10	ug/L	330	ug/Kg
Di-n-octylphthalate	SW-8270B	GC/MS	10	ug/L	330	ug/Kg
Diphenylhydrazine	sw-8270B	GC/MS	10	ug/L	330	ug/Kg
Diphenylamine	sw-8270B	GC/MS	20	ug/L	660	ug/Kg
Disulfoton	SW-8270B	GC/MS	20	ug/L	660	ug/Kg
Ethyl methanesulfonate	SW-8270B	GC/MS	20	ug/L	660	ug/Kg
Famphur	SW-8270B	GC/MS	10	ug/L	330	ug/Kg
Fluoranthene	SW-8270B	GC/MS	10	ug/L	330	ug/Kg
Fluorene	SW-8270B	GC/MS	10	ug/L	330	ug/Kg
Hexachlorobenzene	SW-8270B	GC/MS	10	ug/L	330	ug/Kg
Hexachlorobutadiene	SW-8270B	GC/MS	10	ug/Ł	330	ug/Kg
Hexachlorocyclopentadiene	s₩-8270B	GC/MS	10	ug/L	330	ug/Kg

Parameter	Method Reference	Method Description		Report	ing Limi	t
				eous	Noi	n-Aqueous
Hexachloroethane	SW-8270B	GC/MS	10	ug/L	330	ug/Kg
Hexach lorophene	SW-8270B	GC/MS	500	ug/L	16,500	ug/Kg
Hexachloropropene	SW-8270B	GC/MS	30	ug/L	990	ug/Kg
Indeno(1,2,3-cd)pyrene	SW-8270B	GC/MS	10	ug/L	330	ug/Kg
Isodrin	SW-8270B	GC/MS	30	ug/L	990	ug/Kg
Isophorone	SW-8270B	GC/MS	10	ug/L	330	ug/Kg
Isosafrole	SW-8270B	GC/MS	20	ug/L	660	ug/Kg
Kepone	SW-8270B	GC/MS	250	ug/L	8,250	ug/Kg
Methapryilene	S₩-8270B	GC/MS	100	ug/L	3,300	ug/Kg
3-Methylcholanthrene	SW-8270B	GC/MS	30	ug/L	990	ug/Kg
Methyl methanesulfonate	SW-8270B	GC/MS	20	ug/L	660	ug/Kg
2-Methylnapthalene	sw-8270B	GC/MS	10	ug/L	330	ug/Kg
Methyl parathion	SW-8270B	GC/MS	20	ug/L	660	ug/Kg
Naphthalene	SW-8270B	GC/MS	10	ug/L	330	ug/Kg
1,4 Napthoquinone	SW-8270B	GC/MS	100	ug/L	3,300	ug/Kg
l-Napthylamine	SW-8270B	GC/MS	30	ug/L	990	ug/Kg
2-Napthylamine	SW-8270B	GC/MS	30	ug/L	990	ug/Kg
litrobenzene	SW-8270B	GC/MS	10	ug/L	330	ug/Kg
2-Nitroaniline	SW-8270B	GC/MS	15	ug/L	495	ug/Kg
3-Nitroaniline	SW-8270B	GC/MS	15	ug/L	495	ug/Kg
4-Nitroaniline	SW-8270B	GC/MS	15	ug/L	495	ug/Kg
-Nitroquinoline-1-oxide	SW-8270B	GC/MS	30	ug/L	990	ug/Kg
-Nitrosodi-n-butylamine	SW-8270B	GC/MS	20	ug/L	660	ug/Kg
l-Nitrosodiethylamine	SW-8270B	, GC/MS	30	ug/L	660	ug/Kg
I-Nitrosodimethylamine	SW-8270B	GC/MS	10	ug/L	330	ug/Kg

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Table 9.4. Continued

Parameter	Method Reference	Method Description		Reporti	ng Limi	t
	Reference	bescription ::	Aqu	eous	No	n-Aqueous
N-Nitrosodiphenylamine	sw-8270B	GC/MS	10	ug/L	330	ug/Kg
N-Nitrosodipropylamine	SW-8270B	GC/MS	10	ug/L	330	ug/Kg
N-Nitrosomethylethylamine	SW-8270B	GC/MS	20	ug/L	660	ug/Kg
N-Nitrosomorpholine	sw-8270B	GC/MS	20	ug/L	660	ug/Kg
N-Nitrosopiperidine	SW-8270B	GC/MS	20	ug/L	660	ug/Kg
N-Nitrosopyrrolidine	SW-8270B	GC/MS	20	ug/L	660	ug/Kg
5-Nitro-o-toluidine	SW-8270B	GC/MS	20	ug/L	660	ug/Kg
Parathion	SW-8270B	GC/MS	20	ug/L	660	ug/Kg
Pentachlorobenzene	SW-8270B	GC/MS	20	ug/L	660	ug/Kg
Pentachloronitrobenzene	SW-8270B	GC/MS	20	ug/L	660	ug/Kg
Phenacetin	SW-8270B	GC/MS	20	ug/L	660	ug/Kg
Phenanthrene	SW-8270B	GC/MS	10	ug/L	<b>33</b> 0	ug/Kg
p-Phenylenediamine	SW-8270B	GC/MS	30	ug/L	990	ug/Kg
Phorate	SW-8270B	GC/MS	20	ug/L	660	ug/Kg
2-Picoline	SW-82708	GC/MS	20	ug/L	660	ug/Kg
Pronamide	S₩-8270B	GC/MS	20	ug/L	660	ug/Kg
Pyrene	S₩-8270B	GC/MS	10	ug/L	330	ug/Kg
Pyridine	S₩-8270B	GC/MS	10	ug/L	330	ug/Kg
Safrole	SW-8270B	GC/MS	20	ug/L	660	ug/Kg
Sulfotepp	SW-8270B	GC/MS	20	ug/L	660	ug/Kg
1,2,4,5-Tetrachlorobenzene	SW-8270B	GC/MS	20	ug/L	660	ug/Kg
Thionazin	SW-8270B	GC/MS	20	ug/L	660	ug/Kg
o-Toluidine	SW-8270B	GC/MS	20	ug/L	660	ug/Kg
1,2,4-Trichlorobenzene	SW-8270B	GC/MS	10	ug/L	330	ug/Kg
Triethyl phosphorothicate	SW-8270B	GC/MS	20	ug/L	660	ug/Kg

Parameter	Method Reference	Method Description		Report	ing Limi	t
rarameter		<i>P P P P P P P P P P</i>	Aqu	eous	No.	n-Aqueous
1,3,5-Trinitřobenzene	SW-8270B	GC/MS	30	ug/L	990	ug/Kg
Benzoic Acid	SW-8270B	GC/MS	50	ug/L	1,650	ug/Kg
4-Chloro-3-methylphenol	SW~8270B	GC/MS	_10	ug/L	330	ug/Kg
2-Chlorophenol	SW-8270B	GC/MS	10	ug/L	330	ug/Kg
2,4-Dichlorophenol	SW-8270B	GC/MS	10	ug/L	330	ug/Kg
2,6-Dichlorophenol	SW-8270B	GC/MS	10	ug/L	330	ug/Kg
2,4-Dimethylphenol	SW-8270B	GC/MS	10	ug/L	330	ug/Kg
2,4-Dinitrophenol	SW-8270B	GC/MS	10	ug/L	330	ug/Kg
2-Methyl-4,6-dinitrophenol	sw-8270B	GC/MS	10	ug/L	330	ug/Kg
2-Nitrophenol	sW-8270B	GC/MS	10	ug/L	330	ug/Kg
4-Nitrophenol	SW-8270B	GC/MS	10	ug/L	330	ug/Kg
Pentachlorophenol	sW-8270B	GC/MS	10	ug/L	330	ug/Kg
Phenol	SW-8270B	GC/MS	10	ug/L	330	ug/Kg
2,3,4,6-Tetrachlorophenol	SW-8270B	GC/MS	20	ug/L	660	ug/Kg
2,4,5-Trichlorophenol	sW-8270B	GC/MS	10	ug/L	330	ug/Kg
2,4,6-Trichlorophenol	sW-8270B	GC/MS	10	ug/L	330	ug/Kg
2-Methylphenol	SW-8270B	GC/MS	10	ug/L	330	ug/Kg
3 & 4-Methylphenol	sw-8270B	GC/MS	10	ug/L	330	ug/Kg
sticides/PCBS						
Aldrin	A0808-W2	GC/ECD	0.2	ug/L	500	ug/Kg
Chlordane	SW-8080A	GC/ECD	0.2	ug/L	500	ug/Kg
Dieldrin	A0808-W2	GC/ECD	0.2	ug/L	500	ug/Kg
4,4'-DDD	SW-8080A	' GC/ECD	0.2	ug/L	500	ug/Kg
4,4'-DDE	SW-8080A	GC/ECD	0.2	ug/L	500	ug/Kg

Donomoton	Method Reference	Method: Description	Report	Reporting Limit	
Parameter	Kerer enec	besci iption	Aqueous	Non-Aqueous	
4,4'-DDT	SW-8080A	GC/ECD	0.2 ug/L	500 ug/Kg	
Endosulfan I	A0808-W2	GC/ECD	0.2 ug/L	500 ug/Kg	
Endosulfan II	SW-8080A	GC/ECD	0.2 ug/L	500 ug/Kg	
Endosulfan Sulfate	SW-8080A	GC/ECD	0.2 ug/L	500 ug/Kg	
Endrin	A0808-W2	GC/ECD	0.2 ug/L	500 ug/Kg	
Endrin Aldehyde	SW-8080A	GC/ECD	0.2 ug/L	500 ug/Kg	
Endrin Ketone	SW-8080A	GC/ECD	0.2 ug/L	500 ug/Kg	
Heptachlor	SW-8080A	GC/ECD	0.2 ug/L	500 ug/Kg	
Heptachlor Epoxide	SW-8080A	GC/ECD	0.2 ug/L	500 ug/Kg	
alpha-BHC	SW-8080A	GC/ECD	0.2 ug/L	500 ug/Kg	
beta-BHC	SW-8080A	GC/ECD	0.2 ug/L	500 ug/Kg	
gamma-BHC (Lindane)	SW-8080A	GC/ECD	0.2 ug/L	500 ug/Kg	
delta-BHC	SW-8080A	GC/ECD	0.2 ug/L	500 ug/Kg	
Methoxychlor	SW-8080A	GC/ECD	0.2 ug/L	500 ug/Kg	
Toxaphene	SW-8080A	GC/ECD	0.5 ug/L	500 ug/Kg	
PCB-1016	SW-8080A	GC/ECD	0.2 ug/L	500 ug/Kg	
PCB-1221	SW-8080A	GC/ECD	0.2 ug/L	500 ug/Kg	
PCB-1232	SW-8080A	GC/ECD	0.2 ug/L	500 ug/Kg	
PCB-1242	SW-8080A	GC/ECD	0.2 ug/L	500 ug/Kg	
PCB-1248	A0808-W2	GC/ECD	0.2 ug/L	500 ug/Kg	
PCB-1254	SW-8080A	GC/ECD	0.2 ug/L	500 ug/Kg	
PCB-1260	SW-8080A	GC/ECD	0.2 ug/L	500 ug/Kg	
otal Petroleum Hydrocarbons	EPA 418.1	1R	2.0 mg/L	10 mg/Kg	
otal Petroleum Hydrocarbons (Diesel Range Organics)	SW-8015B	GC	0.1 mg/L	4.0 mg/Kg	

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Parameter	Method Reference	Method Description	İ	Reportin	g Limit	
rai directes	Kerel elle		Aqueous		Non-Aqueous	
Total Petroleum Hydrocarbons (Gasoline Range Organics)	SW-8015A Modified/SW-8015B	GC	0.1	mg/L	0.5 mg	ı/Kg
HPLC - Polynuclear Aromatic Hydro	ocarbons					
Napthalene	SW-8310	HPLC	2.0	ug/L	200	ug/Kg
Acenaphthylene	SW-8310	HPLC	1.0	ug/L	100	ug/Kg
Acenaphthene	SW-8310	HPLC	1.0	ug/L	100	ug/Kg
Fluorene	SW-8310	HPLC	1.0	ug/L	100	ug/Kg
Phenanthrene	SW-8310	HPLC	1.0	ug/L	100	ug/Kg
Anthracene	SW-8310	HPLC	2.0	ug/L	100	ug/Kg
Fluoranthene	SW-8310	HPLC	0.2	ug/L	20	ug/Kg
Pyrene	SW-8310	HPLC	0.2	ug/L	20	ug/Kg
Benzo(a)anthracene	SW-8310	HPLC	0.2	ug/L	20	ug/Kg
Chrysene	SW-8310	HPLC	0.2	ug/L	20	ug/Kg
Benzo(b)fluoranthene	sw-8310	HPLC	0.2	ug/L	20	ug/Kg
Benzo(k)fluoranthene	SW-8310	HPLC	0.2	ug/L	20	ug/Kg
Benzo(a)pyrene	SW-8310	HPLC	0.2	ug/L	20	ug/Kg
Dibenz(ah)anthracene	SW-8310	HPLC	0.2	ug/L	20	ug/Kg
Benzo(ghi)perylene	S₩-8310	HPLC	0.2	ug/L	20	ug/Kg
Indeno(1,2,3-cd)pyrene	sw-8310	HPLC	0.2	ug/L	20	ug/Kg

25° 44 Table 9.5. Analytical Methods and Reporting Limits - Misc.

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Parameter	Method Reference	Method Description	Ro	eporting Li	g Limít nît
		<b>*</b>	Aqueo	us	Non-Aqueous
Acidity	EPA-305.2	Titration	10. r	mg/L	NA
1,2-Dibromo-3-chloropropane	EPA-504.1	GC	0.02 (	ug/L	NA
Ethylene dibromide	EPA-504.1	GC	0.02	ug/L	NA

## 10. DATA REDUCTION, VALIDATION AND REPORTING

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#### 10.1 Data Reduction

All analytical data are reduced to the appropriate concentration units as specified by the method. The analyst will reduce the data taking into account any and all factors such as dilution, percent solid, sample weight or volume and reagent normality. Blank correction will be applied only when required by the method.

## 10.2 Data Validation

Data validation is the process by which data is accepted or rejected based on pre-determined criteria. TestAmerica does not provide data validation services.

### 10.3 Data Verification

- 10.3.1. Data is evaluated based on the following broad range of criteria:
  - Proper sample collection, storage and holding time.
  - Use of standard operating procedures or other approved analytical procedures.
  - Use of properly operating and calibrated instruments.
  - Successful analysis of appropriate quality indicators.
- 10.3.2. All data will be evaluated and verified prior to being released for reporting purposes to the TestAmerica Project Management team. The persons evaluating the data will have sufficient knowledge of the technical work to identify questionable values. All raw data and pertinent record are maintained for a period of 7 years for non-potable data and 10 years for potable data. as part of the Voluntary Action Program (VAP) requirements, all documents prepared or acquired in connection with a voluntary action will be retained for a period of 10 years from the date the analyses were submitted to a certified professional.
- All analytical data will be verified for completeness of Quality Control Indicator requirements, and will be spot checked for completeness. This verification will be performed by a competent analyst or the area supervisor.

After an analyst completes training on a parameter, and passes a PE sample, he/she will be permitted to perform self verification of data using specific forms designed for this purpose.

Data which is determined to be of questionable quality, either due to reasons initiating from the laboratory or concerns voiced by the client, will be reviewed by a member of the laboratory management staff. Clients will be informed of any and all data which does not meet the full Quality Control requirements as outlined in the various standard operating procedures.

10.3.3. The laboratory will use the Intra-Laboratory Notification form, Figure 10.2, to communicate any quality issues or special circumstances (i.e. especially bad matrix, holding time issues, etc.) to various members of the laboratory. The Re-Evaluation Request form, Figure 10.3, is used by the Project Managers to request a re-evaluation of a sample, describing the required action(s) to take and the sample number(s)) in question. Response information such as the reason for the difference noted, problem corrected, and the type of subsequent action necessary is collected. These two forms are retained in respective project files.

The Intra-Laboratory Notification form is also used by members of the laboratory staff to communicate either internal complaints, or complaints from customers, to members of the management in order that they may be examined and resolved.

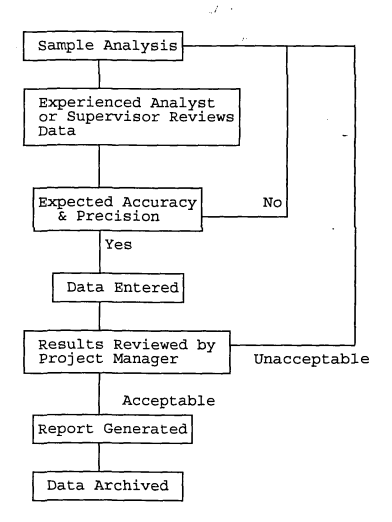
## 10.4 Data Reporting

Analytical results will be reported in a manner acceptable to the client. All reports will be assembled and approved by the Project Management team and delivered to the client within the time period agreed upon by the client and the laboratory. Data is generally reported at the limit of quantitation (LOQ) (Reporting Limit = LOQ). The LOQ is determined for most analytes by performing a method detection limit (MDL) study. The protocol used to determine the MDL is found in 40 CFR Part 136 Appendix B. Analytical methods and reporting limits for analytes are listed in the Tables in Section 9.

Additional data required by the customer, such as operating conditions, quality control data, method detection limits (MDLs), recommendations or problems will be reported by the Project Management team.

Figure 10.1 shows the analytical data review and reporting scheme utilized by TestAmerica-Dayton.

Figure 10.1. Analytical Data Review and Reporting Scheme



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# Figure 10.2. Intra-Laboratory Notification Form INTRA-LABORATORY NOTIFICATION FORM

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DATE INITIATED:	A 1
SAMPLE NUMBER(S):	, 
PARAMETER:	DEPARTMENT:
CLIENT:	SUPERVISOR:
DEVIATION/CONCERN:	
ACTION RECOMMENDED:	
CLIENT CONTACT YES	NO
CONTACT NAME	DATE:
COMMENTS:	
PROJECT MANAGER LABORATORY MANAGER DIVISION MANAGER OA/OC COORDINATOR	

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# Figure 10.3. Re-Evaluation Request Form RE-EVALUATION REQUEST FORM

		-	
DEPARTMENT:		JOB NUMBER:	
PARAMET	TER:	CLIENT:	
DUE DAT	TE:		
DATE CO	OMPLETE:	REQUE	ST DATE:
SAMPLE I.D.	ORIGINAL RESULT	RER RESULT	EXPLANATION
ACTION REQU	JESTED:	<del>-</del>	CHECK DATA ENTRY
	CHECK CALCULATION	<del></del>	REPEAT ANALYSIS
	CHECK QC		OTHER
REASON FOR	REQUEST:		
		<del></del>	
ACTION TAKE	en:	ROUTING:	
	CONTACTED CLIENT		DEPT. SUPERVISOR
	NO ACTION NEEDED		DATA REVIEW AND APPROVAL
	ENTERED NEW RESULTS		PROJECT MANAGER
-	ISSUED CORRECTED REP	ORT	QA/QC OFFICER
	OTHER		LAB MANAGER

#### 11. INTERNAL QUALITY CONTROL AND QUALITY ASSURANCE

## 11.1 Internal Quality Control

Internal quality control makes use of several types of QC samples to monitor the performance of the measurement process. Quality control checks are analyzed to ensure the generation of accurate and valid data on client samples. Please refer to Section 5 for control limits for the following QC samples. For information concerning preparation, storage and shelf life of the various Quality Control Indicators (QCI), please refer to the specific parameter SOP.

## 11.1.1 Blank Samples

Blank samples are analyzed to assess the extent (if any) of contamination due to the method, transit or storage. Blank samples related to field sampling are defined in Section 6. These blanks will be supplied by TestAmerica based on the data quality objectives of the project.

Blank samples which are performed with analyses include:

Method Blank The method blank is prepared just like a sample. The method blank is analyzed with samples which are processed at the same time as the blank to assess the extent of contamination obtained during the preparation process.

Solvent/Reagent Blank The reagent blank is prepared from the same lot of solvent or reagent used in the analysis. It is used to assess the background of solvents/reagents.

### 11.1.2 Surrogate Compounds

Surrogates are known concentrations of compounds which are added to every blank, sample, matrix spike, matrix spike duplicate and standard in order to evaluate the analytical efficiency of the method in individual sample matrices. The surrogate compounds are chemically similar to the target compounds. Surrogates are utilized based on method requirements.

#### 11.1.3 Calibration Verification

Verification samples are analyzed during each run to assure that the method and/or instrument is properly calibrated and that calibration is maintained throughout the analytical run. Calibration verification standards include:

Initial Calibration Verification (ICV) A standard which is analyzed from a source different from those used for calibration to check the validity of the initial calibration curve. If the ICV does not pass QC criteria, the ICV is re-analyzed. If the ICV still fails QC criteria, analysis is ended, the problem is investigated, and the instrument is re-calibrated. If an ICV is used

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in place of a CCV, it must meet or exceed the quality control requirements of the CCV.

Continuing Calibration Verification (CCV) A standard which is analyzed during the analytical run to confirm calibration. CCVs must meet the quality control requirements listed within the specific method. All client samples must be bracketed by acceptable CCVs.

#### 11.1.4 Internal Standards

Internal standards are compounds which are added to every standard, blank, matrix spike, matrix spike duplicate, and sample at a known concentration prior to the analysis. The internal standards are used as the basis for quantitation of the target compounds. The utilization and recovery of the internal standards must meet method-specific guidelines. If control limits cannot be met, the sample(s) are re-analyzed. If samples cannot be re-analyzed due to limited sample volume or holding time issues, the results are flagged and the client is notified.

#### 11.1.5 Spiked Samples

The laboratory analyzes samples which have been fortified, or spiked, with known concentrations of target analytes. Spiked samples are analyzed for a variety of reasons, and include:

Matrix Spike/Matrix Duplicate (MS/MSD) Two aliquots of sample are spiked with the analyte(s) and the recovery is determined. The matrix spike (MS) recovery indicates the presence or absence of matrix interferences, and the duplicate sample analysis (MSD) is carried out to verify precision.

Analytical Spike (AS) An aliquot of digested sample or sample into which a known amount of compound is added. The analytical spike is analyzed immediately and the recovery is calculated in order to assess the matrix effect on the analytical system.

Laboratory Control Sample (LCS) A control sample of known composition. Control samples are analyzed using the same sample preparation reagents and analytical methods as employed for samples in order to verify that the preparation and analysis methods are in control.

#### 11.1.6 Duplicate Samples

A duplicate sample is a second aliquot of a sample which is carried through sample preparation and analysis procedures to verify the precision of the analytical method for that matrix.

## 11.2 Reagent and Standards Quality Control

Reagents used in the laboratory are of analytical reagent grade or higher purity. Reagent lots are checked by the analysis of reagent blanks. A reagent is labeled at the time of receipt with the date received, who received it, expiration date, manufacturer's lot number and date opened.

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Material Safety Data (MSDS) are on file for all hazardous chemicals and available to all analysts. Reagents are stored in a designated reagent storage area. As appropriate, smaller quantities are stored in ventilated solvent cabinets in the laboratories and in accordance with the Material Safety Data Sheet (MSDS) storage requirements.

Records are maintained for all standards. All standards are logged into the appropriate standards logbooks which contain records of manufacturer, expiration of the standard and concentration (or purity).

### 11.3 Performance Evaluation Samples

Standard Reference Materials (SRMs) or any other appropriate known concentrates are analyzed on a routine basis as a quality control check. These samples are analyzed along with regular samples in the normal laboratory routine. The analyst compares the results with the known values and with the acceptance criteria outlined in Section 5. Performance Evaluation (PE) samples are utilized to document analyst training and to verify that analytical systems remain in control.

## 11.4 Internal Quality Assurance

To monitor quality, the following actions are periodically taken by the Division/Operations Manager(s) and Quality Assurance Coordinator:

## 11.4.1 QC Single or Double Blind Samples

Samples which are known to be PE samples (single blind) and samples which are not known to be PE samples (double blind) are prepared by the QA Coordinator on a periodic basis or when requested by Division/Project Manager(s) to assess analysis. These samples are analyzed and the results are reported to the divisional QA Coordinator. The QA Coordinator then reviews the analytical data and determines if corrective action is needed.

#### 11.4.2 Internal Audits

Periodically, internal audits are conducted by the divisional QA Coordinator to evaluate systems and performance as described in Section 12.

#### 12. SYSTEM AND PERFORMANCE AUDITS

12.1 Performance Audits

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Performance audits provide a systematic check of laboratory data quality and measurement systems. For maximum usefulness two types of performance evaluation samples are employed, single blind and double blind.

Single-blind A sample which is known by all concerned to be a PE sample and only the values are unknown. The results of these samples are useful in determining technical systematic problems within the operating group.

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Double-blind A sample that appears to be a client sample; its identity and values are both unknown to the laboratory. Double-blind samples are useful in identifying technical systematic problems, random analytical problems, and non-technical systematic problems.

TestAmerica Dayton routinely participates in single-blind laboratory performance evaluations.

#### 12.2 Systems Audits

A system audit is an evaluation of a laboratory's quality assurance practices and operating procedures. This audit consists of an on-site review of the laboratory's quality assurance systems and its physical facilities. In addition to internal audits performed by the QA Coordinator, periodic systems audits are performed by the Director of Data Quality. Findings of these audits are reported in writing to the Division Manager and the Corporate Office. If appropriate, corrective action is requested and the corrective action taken is documented. Clients and regulatory agencies may also perform system audits.

- 12.2.1 The system audit may include any of the following:
  - Personnel, facilities and equipment;
  - Chain-of-custody procedures;
  - Sample tracking procedures;
  - Instrument calibration and maintenance;
  - Standards preparation and verification;
  - Sample preparation procedures;
  - Analytical procedures;
  - Quality Control procedures;

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- Data handling procedures;
- Training records;
- Documentation; and
- Document control procedures.

#### 13. PREVENTATIVE MAINTENANCE PROCEDURES AND SCHEDULES

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### 13.1 Preventative Maintenance Program

TestAmerica follows a well-defined program to prevent the failure of laboratory equipment or instrumentation during use. This program of preventative maintenance helps to avoid delays due to instrument downtime. Adequate supplies of spare parts such as GC columns, syringes, septa, injection port liners and electronic parts are maintained in the laboratory.

Routine preventative maintenance procedures such as lubrication, source cleaning, detector cleaning and the frequency of such maintenance are performed according to the procedures outlined in the manufacturer's manual. Chromatographic carrier gas purification traps, injection port liners and septa are cleaned or replaced on a regular basis. Precision and accuracy data are examined for trends and excursions beyond established control limits to determine evidence of instrument malfunction. Maintenance must be performed by laboratory analysts when there is evidence of degradation of peak resolution, a shift in the calibration curves, loss of sensitivity, or failure to meet one of the quality control criteria.

The preventative maintenance performed on major laboratory instrumentation is summarized in Table 13.1. Instrument logbooks containing usage, calibration, maintenance and repair records are kept in the laboratories at all times.

## 13.2 Equipment Malfunction

In the event of equipment malfunction that cannot be resolved within two working days, service shall be obtained from the instrument vendor or manufacturer, if such a service agreement exists or can be tendered. If on-site service in the laboratory is unavailable, arrangements shall be expedited to have the instrument shipped to the manufacturer for repair. Back-up instruments which have been approved for the analysis shall perform the analysis normally carried out by the malfunctioning instrument, if feasible. If back-up is not available and the analysis cannot be carried out within the needed time frame, the samples shall be subcontracted to another approved laboratory to carry out the analysis.

Table 13.1. Maintenance Procedures for Major Instrumentation

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Instrumentation	Maintenance Procedure	Spare Parts
Instrumentation	Maintenance Procedure	Spare Faits
Gas Chromatograph/Mass Spectrometer	<ol> <li>Replace pump oil as needed.</li> <li>Change septa as needed.</li> <li>Change gas line dryers as needed.</li> <li>Clean source as needed.</li> <li>Replace electron multiplier as needed.</li> <li>Injection port cleaning as needed.</li> </ol>	Syringe Septa Various electronic components Plumbing supplies Injection port liners
Gas Chromatograph	<ol> <li>Change septa as needed.</li> <li>Clean gas line dryers as needed.</li> <li>Change syringes on autosamplers as needed.</li> <li>Leak check when installing columns</li> <li>Injection port cleaning as needed.</li> <li>Check inlet system for residue buildup periodically.</li> </ol>	Syringe Septa Various electronic components Plumbing supplies Injection port liners
Purge and Trap Sample Concentrator	<ol> <li>Replace trap as needed.</li> <li>Decontaminate system as required by blank analysis.</li> <li>Check system for leaks.</li> </ol>	Traps Various electronic components Plumbing supplies
Graphite Furnace Atomic Absorption Spectrophotometer	<ol> <li>Change graphite contact rings as needed.</li> <li>Clean quartz windows as needed.</li> <li>Change tubes as needed.</li> </ol>	Contact rings Tubes

Table 13.1 Continued...

Instrumentation	Maintenance Procedure	Spare Parts
Inductively Coupled Plasma Spectrometer	<ol> <li>Change sample rinse lines.</li> <li>Clean nebulizer components, torch assembly and spray chamber.</li> <li>Clean filters.</li> <li>Clean mirrors.</li> </ol>	Nebulizer components Torch assembly Pump tubing and sample probe
Inductively Coupled Plasma - Mass Spectrometer	<ol> <li>Change pump tubing as needed.</li> <li>Clean nebulizer components, torch assembly and spray chamber.</li> <li>Clean sampler and skimmer cones.</li> <li>Change roughing pump oil.</li> </ol>	Spare electrode Pump tubing Nebulizer components Torch assembly Spray chamber assembly Sampler and skimmer cones Pump oil
pH/Conductivity Meter	<ol> <li>Clean electrodes as needed.</li> <li>Refill electrodes as needed.</li> </ol>	Filling solution
Balance	<ol> <li>Check level of balance daily.</li> <li>Clean balance pan daily.</li> <li>Weigh and record a known mass daily.</li> <li>Calibrate and clean balance monthly.</li> <li>Outside service on all balances annually.</li> </ol>	
Wet Chemistry Auto Analyzer	<ol> <li>Recharge/replace coils as needed.</li> <li>Clean/replace flow cells as needed.</li> <li>Change pump tubes and gas line as needed.</li> <li>Clean sampling pivot head and replace probe as needed.</li> </ol>	Glass connectors Tubing Glass coils (5 and 20 turn) Cd reduction coils

Table 13.1. Continued...

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Instrumentation	Maintenance Procedure	Spare Parts
Total Organic Carbon Analyzer (TOC)	<ol> <li>Change copper/tin scrubber as needed.</li> <li>Clean combustion tube as needed.</li> <li>Replace permeation dryer when discolored.</li> <li>Check and clean IC chamber, TC inlet valve, IC inlet valve, bottom connector and ASM sample loop as needed.</li> </ol>	Septa Sample tip Copper/tin particles
Mercury Analyzer	<ol> <li>Change drying tube daily.</li> <li>Change pump tubing weekly.</li> <li>Clean optical cell as needed.</li> <li>Clean liquid/gas separator as needed.</li> </ol>	Assorted Tubing Hg Lamp Liquid/Gas Separator Assembly

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## 14. SPECIFIC ROUTINE PROCEDURES TO ASSESS DATA PRECISION, ACCURACY AND COMPLETENESS, AND OTHER QUALITY CONTROL INDICATORS

### 14.1 Precision

Precision is a measure of the degree of agreement between repeated measurements of the same parameter under prescribed, similar conditions. Analytical precision will be monitored using results from duplicate analyses. Analytical precision goals expressed as relative percent difference (RPD), are presented in Section 5. The RPD is calculated as follows:

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$$RPD = \underline{Absolute\ Value\ (D1) - (D2)} \times 100$$

$$(D1 + D2)/2$$

where,

RPD is the relative percent difference

D1 is the first duplicate value (percent recovery); and

D2 is the second duplicate value (percent recovery).

#### 14.2 Accuracy

Accuracy is a measure of the degree of agreement between an analyzed value and the true or accepted reference value. The accuracy of a measured value is expressed as a percent of the expected or known value. In the laboratory, accuracy will be evaluated by comparing the recoveries of parameters of interest against criteria outlined in Section 5, through the use of quality control reference samples or reference materials. The recovery of a compound will be defined as:

$$%R = \frac{(SSR - SR)}{S} \times 100$$

where,

%R is percent recovery
SSR is the spiked sample result
SR is the sample result; and
S is the spike concentration

## 14.3 <u>Completeness</u>

Completeness is a measure of the amount of valid data obtained from the samples received. It is defined in terms of a percentage of the number of valid measurements expected. Ideally, every sample will generate all of the valid measurements expected. Realistically, some samples may be lost in laboratory accidents or some data may be deemed questionable based on internal quality control criteria. Such instances will be documented and communicated to the client in a narrative section of the report.

Completeness also implies the ability of the final report to answer the client's questions. TestAmerica will have personnel available to discuss analytical reports with clients. Every attempt will be made by TestAmerica to achieve 100% completeness on analytical parameters. All judgements of completeness will be determined by the client. Percent completeness is calculated as follows:

where,

% C = Percent Completeness

V = number of results judged to be valid

n = total number of results

### 14.4. Other Quality Control Indicators (QCI)

## 14.4.1. Method Detection Limit (MDL) Studies

Method Detection Limit Studies are calculated using between seven and ten replicates. The equation is as follows:

MDL = SD \* Student's T Value

where,

SD = the Standard Deviation of the seven to ten replicates

Student's T Value = value based on the number of
 replicates (see below):

Student's T Value
2.821
2.896
2.998
3.143

For additional information, please refer to the SOP for detection limit studies.

14.4.2. Statistically Based Control Limits

14.4.2.1. Statistically based control limits are calculated using a minimum of twenty data points. The individual limits are calculated as follows:

Upper Control Limit (UCL) = Mean + 3SD Upper Warning Limit (UWL) = Mean + 2SD Lower Warning Limit (LWL) = Mean - 2SD Upper Control Limit (LCL) = Mean - 3SD

where,

ITEM OR TREND

2 consecutive points within

warning limits

Mean = the Average of the replicates SD = The Standard Deviation of the replicates

POSSIBLE INDICATION

Procedure is out of

must be corrected

control and the problem

14.4.2.2. Trend Analysis, using the control limits, is a useful tool in helping to identify when a procedure is out of control or approaching an out of control situation. Some items or trends to look for and what they may indicate are as follows:

Any point outside of the control limits	Out of control
7 consecutive points increasing or decreasing	Approaching out of control situation
Cycles or reoccurring patterns	There is a variable in the procedure that is affecting results
7 data points on the same side of the center line	Something in the procedure has changed and is affecting results

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#### 15. CORRECTIVE ACTION

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An important part of any quality assurance program is a well-defined, effective policy for correcting quality problems. NET maintains a corrective action system which operates under the direction of the Division Manager and Quality Assurance Coordinator. While the entire quality assurance program is designed to avoid problems, it also serves to identify and correct those that occur. Usually these quality problems fall into two categories: immediate corrective action or long-term corrective action.

## 15.1 Immediate Corrective Action

Specific quality control procedures are designed to help analysts detect the need for corrective action. Often, an analyst's experience will be most valuable in identifying abnormal analyses or malfunctioning equipment. Immediate corrective action may be taken. Such actions should be noted in laboratory notebooks but no other formal documentation is required unless the corrective action taken fails to correct the problem.

## 15.2 Long Term Corrective Action

The need for formal corrective action may be identified by performance on routine QC samples, control chart trends, or as a result of a performance or systems audit. Any quality problem which cannot be solved by immediate corrective action falls into this category. The division QA Coordinator is responsible for managing the corrective action process and communicating the status of corrective action progress to the Division Manager.

The QA Coordinator may, with the support of the Division Manager, delegate responsibilities for investigating problems and implementing solutions to appropriate operational groups or individuals. Involvement of the analyst and supervisor of the area concerned is crucial to the effectiveness of the corrective action process. It is the responsibility of analysts and supervisors to write corrective action reports, and it is the responsibility of the QA Coordinator to maintain the corrective action reports.

- 15.2.1. The essential steps in the closed loop corrective action system are:
  - 1. Identification of the problem
  - 2. Assignment of responsibility for investigating the problem
  - 3. Determination of the cause of the problem through investigation

- 4. Formulation of a corrective action plan
- 5. Assignment of responsibility for implementation of the corrective action plan
- 6. Monitoring the effectiveness of the corrective action plan
- 7. Verifying the elimination of the problem
- 8. Documenting the process involved

## 15.3 Corrective Action Reports

Corrective Action Reports are formal documentation of long term corrective action taken at the Division. These reports are required for "Unacceptable" results on Performance Evaluation (PE) studies.

- 15.3.1 Steps Required to Complete a Corrective Action Report
  - 1. Notification of acceptability of results.
  - 2. Quality Assurance Coordinator informs appropriate analyst, supervisor, and Project Managers of unacceptable parameters requiring a Corrective Action Report (CAR).
  - 3. Analyst determines, through careful and thorough consideration, possible sources of the problem.
  - 4. Analyst with the help of the Supervisor, if necessary, identifies the assignable cause of the problem and documents this on the CAR form.
  - 5. Along with identification of the problem, the specific steps taken to correct the problem are documented on the CAR form.
  - 6. The analyst reviews the CAR with the Supervisor, the QAC, the Division Manager and/or the Project Manager to ensure that the assignable cause is understood and agreed upon.
  - 7. If appropriate, after the problem has been identified and corrected, a blind performance evaluation sample is submitted by the QA Coordinator.
  - 8. Successful completion of the blind performance sample will demonstrate that the analysis is in control. Unsuccessful completion of the blind performance sample will indicate that appropriate corrective action has not taken place and the process must start over with analyst identification of the problem.
  - 9. After successful completion of the corrective action

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process, the CAR is reviewed and signed by the QA Coordinator and the Division Manager.

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The QA Coordinator verifies corrective action is maintained in the laboratory by reviewing analytes with CARs during his/her routine systems audits.

#### 16.0 QUALITY ASSURANCE REPORTS TO MANAGEMENT

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## 16.1 Quality Assurance Coordinator - Reports to Management

The Quality Assurance Coordinator is responsible for reporting to management on the effectiveness of the Quality Assurance Plan. A monthly summary of quality-related issues is prepared and submitted to the Director of Data Quality, the Division Manager, and other appropriate personnel.

- 16.2.1. The monthly quality assurance report topics are:
  - A. Any Key Issues
  - B. SOPs
  - C. Corrective Action Reports
  - D. MDLs
  - E. Audits and Client Visits
  - F. Performance Evaluation Samples
  - G. Certification, Accreditation and Contract Approval
  - H. Training
  - G. Other

#### 16.2 Quality Systems Management Review

It is our policy for the senior divisional management team to conduct an annual review of its quality systems to ensure its continuing suitability and effectiveness in meeting client and regulatory requirements and to introduce any necessary changes or improvements.

This review uses information generated during the preceding year to assess the "big picture" by ensuring that routine quality actions taken and reviewed on a monthly basis are not components of larger systematic concerns. The monthly review should continually keep the quality systems current and effective, therefore, the annual review is a formal senior management process to review specific existing documentation.

- 16.3.1. The significant issues from the following documentation should be summarized by the Quality Assurance Coordinator prior to the review meeting:
- matters arising from the previous annual review;
- prior monthly Quality Assurance Reports, including information on:

- \_ internal systems audit summaries and corrective actions;
  - reports from audits by clients or third-party assessments;
  - results of performance evaluation samples, including corrective actions implemented;
  - results of internal quality checks;
  - certification / accreditation issues;
  - methods or SOP issues;
  - staff training;
  - prior Re-Evaluation Request forms;
- minutes from prior data quality management and staff meetings;
- minutes from prior Senior Management Team meetings, including:
  - adequacy of staff, equipment and facility resources;
  - future plans for resources and testing capability and capacity;
- prior Customer Service / Business Development meeting information and prior Inter -Laboratory notification forms that involves data quality issues or client complaints.

The annual review includes the previous 12 months and can occur anytime during the calendar year to best meet the needs of the division. Based on the annual review, a report is generated by the Quality Assurance Coordinator for distribution to the Division's Senior Management Team and the Director of Data Quality that includes:

- when the review occurred and who participated;
- a reference to the existing data quality related documents and topics that were reviewed;
- what quality systems changes or improvements will be made as a results of the review;
- an implementation schedule for the changes.

The Divisional Quality Assurance Plan should be revised at this time to reflect any significant changes made to the quality systems.

TestAmerica, Inc. Dayton Division Quality Assurance Plan

## APPENDIX 1. ANALYTICAL EQUIPMENT LIST

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The Dayton Division of TestAmerica maintains a full range of modern, state-of-the-art equipment and instrumentation. Additional equipment and instrumentation is available at other TestAmerica laboratories located throughout the United States.

Listings of major analytical instrumentation and equipment for both the laboratory and field operations are found in Tables 1 - 5 of this Appendix.

Table 1. Equipment List for Metals Department

CAR

: Graphite Furnace Group

- Atomic Absorption unit: Perkin Elmer SIMA 6000

Simultaneous graphite furnace with Zeeman correction

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and autosampler

- Data System: Dell Optiplex GS

- Printer: HP Laserjet 4

: Inductively Coupled Plasma Spectrometer (ICP) Group

- ICP Spectrometer: TJA Model 36

- Autosampler: Model AS 300

- Data System: NPC 486, ThermoSPEC software

- Printer: Epson LQ 570+

: Automated Cold Vapor (Mercury) Group

- Leeman Labs Model PS200 with autosampler

- Printer: GSX-190 Citizen

- Data System: IBM PC with Leeman PS200 software

: Inductively Coupled Plasma - Mass Spectrometer (ICP-MS)

- ICP-MS: Perkin-Elmer ELAN 6000 - Autosampler: Perkin-Elmer AS 91 - Data System: Dell Optiplex 6xi - Printer: HP Laserjet 4+

Unit : Metal Preparation Laboratory

Balance: Mettler AG 204

Table 2. Equipment List for Wet Chemistry Department

Description	Manufacturer	Model
Auto Analyzer with (2) Autosampler	Bran-Luebbe	Traacs 800
TOC Analyzer with Autosampler	Tekmar/Dohrmann	DC-190
Spectrophotometer	Milton Roy	301
Spectrophotometer	Milton Roy	501
Ion Analyzer	Orion Research	901
pH Meter	Orion Research	SA 520
pH Meter	Accumet	20
pH Meter	Orion Research	601 A
Turbidimeter	Hach	2100 AN
Flash Point Analyzer	Precision Scientific	
Oxygen Meter	YSI Scientific	5000
ZHE Extractors	Millipore	
Balance	Mettler	нк 160
Balance	Mettler	AE 160
Muffle Furnace	Lindberg	51828
Vacuum Oven	Fisher Scientific	281
Orbital Shaker	Labline	3590
Rapidstill II	Labconco	
Midi CN Distillers (2)	Labcrest	

Table 2. Continued...

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Description	Manufacturer	Model
Ovens (4)	Blue M / VWR	
Incubators (3)	Precision/Unitherm/Puff	er Hubbard
Water Bath	Precision	180
RADIOLOGY		_
Alpha/Beta/Gamma System with Autosampler (3) and NaI Detector (3)	Canberra	2404
BACTERIOLOGY		
Autoclave	Amsco	57 CR
Microscope	American Optical	110
Colony Counter	American Optical	3352
Incubator	<b>VW</b> R	3020
Water Bath	Blue M	MW 1120A-1
Bacti-Cinerator II	Scientific Products	Cat. No B9753

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Unit: Balance: Mettler BB 330

Table 3. Equipment List for GC/MS Department

-Unit: GC/MS-A (Volatiles) - GC- Hewlett Packard Model 5890 - MS- Hewlett Packard Model 5970 - Liquid Sample Concentrator: Tekmar Model LSC-3000 - Autosampler: Archon Model 5100 - Data System: Hewlett Packard Vectra PC with Enviroquant software - Printer: Hewlett Packerd 4si Unit: GC/MS-B (Semi-Volatiles) - MS- Hewlett Packard Model 5970 - Autosampler: Hewlett Packard Model 7673A - Data System: Hewlett Packard Vectra PC with Enviroquant software Unit: GC/MS-C (Volatiles) - GC- Hewlett Packard Model 5890 - MS- Hewlett Packard Model 5970A - Liquid Sample Concentrator: Tekmar Model LSC-3000 - Autosampler: Archon Model 5100 - Data System: Hewlett Packard Vectra PC with Enviroquant software - Hewlett Packard 4si GC/MS-D (Volatiles) Unit: - GC- Hewlett Packard Model 5890 - MS- Hewlett Packard Model 5970 - Liquid Sample Concentrator: Tekmar Model LSC 2000 - Autosampler: Tekmar Model ALS 2050 - Data System: Hewlett Packard Vectra PC with Enviroquant software Unit: GC/MS-E (Semi-Volatiles) - GC- Hewlett Packard Model 5890 Series II - MS- Hewlett Packard Model 5970 - Autosampler: Hewlett Packard Model 7673A - Data System: Hewlett Packard Vectra PC with Enviroquant software - Hewlett Packard 4si Unit: GC/MS-F (Semi-Volatiles) - GC- Hewlett Packard Model 5890 - MS- Hewlett Packard Model 5970 - Autosampler: Hewlett Packard Model 7673A System: Hewlett Packard Vectra PC with - Data Enviroquant software - Hewlett Packard 4si

## Table 4. Equipment List for GC Department

Unit: GC-1 - GC- Hewlett Packard Model 5880A - Autosampler: Hewlett Packard Model 7671A - Detector: Flame Ionization - Integrator(2): Hewlett Packard Unit: GC-2 - GC- Hewlett Packard Model 5890 - Autosampler : Hewlett Packard Model 7673A - Detector 1 - ECD - Detector 2 - ECD - Data System: Hewlett Packard Vectra PC with Enviroquant Software Unit: GC-3 - GC - Hewlett Packard Model 5890, Series II - Detector - Dual ECDs - Data System: Hewlett Packard Vectra PC with Enviroquant Software - Printer: Hewlett Packard Unit: GC-4 - GC - Hewlett Packard Model 5890, Series II - Detector - Dual PIDs, FID - Autosampler - Tekmar 2000 Conc., Archon autosampler - Data System: Hewlett Packard Vectra PC with Enviroquant Software - Printer: Hewlett Packard Unit: GC-5 - GC- Hewlett Packard Model 5890 - Autosampler : Hewlett Packard Model 7673A - Detector 1 - ECD - Detector 2 - ECD - Data System: Hewlett Packard Vectra PC with Enviroquant Software - Printer: Hewlett Packard IIIsi Unit: GC-6 - GC- Hewlett Packard Model 5890 - Autosampler (2): Hewlett Packard Model 7673A - Detector 1 - ECD - Detector 2 - ECD - Data System: Hewlett Packard Vectra PC with Enviroquant Software

- Printer: Hewlett Packard IIIsi

TestAmerica, Inc. Dayton Division Quality Assurance Plan

## Table 4. Equipment List for GC Department, continued

Unit: GC-7

- GC- Hewlett Packard Model 5890

- Auto Sampler (2): Hewlett Packard Model 7673A

- Detector: Dual PIDs; Trimetrics - Detector: Dual FIDs; Hewlett Packard - Data System: Hewlett Packard Packard Vectra PC with

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Enviroquant Software

- Printer: Hewlett Packard IIIsi

Unit:

- Liquid Chromatography Unit: Hewlett Packard Model 1050

- Flourescence Detector : Hewlett Packard 1046A

- Data System: Hewlett Packard Vectra PC with

Chemstation Software

- Printer: Hewlett Packard II

Unit: FT-IR

- FT-IR: Perkin Elmer 1600 Series - Printer: Okidata Microline 391

Unit: Balance

- Mettler Model AE 163 - Mettler Model PE 360

Unit: Nitrogen sample concentration unit (2)

- Labconco Rapidvap

Table 5. Equipment List for Field Services Department

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Description	Manufacturer	Model
Autosamplers (7)	ISCO	2700
Autosamplers (2)	ISCO	3700
Autosamplers (1)	ISCO	2100
Flow Meters (4)	ISCO	1870
Flow Meters (1)	ISCO	3210
Flow Meter (3)	ISCO	3230
Conductivity Meter	YSI	33
pH Meter (2)	Hanna	9025
Residual Chlorine Kit	Hach	CN66
Water Level Indicator	SINCO	51405301
Pressure Filtration Device	Geotech	0856
Confined Space Entry Equipment		
Ventilator Winch Gas Monitors	Air Systems International Miller Equipment Industrial Scientific	SVB-G8 50 G HMX 271
Field Sampling Vehicles	(4) Chevy and GMC	
4 Inch Well Pump	Suburban	P051-2W
2 Inch Well Pump	Grandfos	Rediflo 2
Electrical Generator	Pincor	RF-30HM5
Power Auger	Tecumseh Engines	21
2 Inch Teflon Bailer	Modern Industrial Plastics	GWE-300